| Ref<br># | Hits | Search Query  | DBs                | Default<br>Operator  | Plurals | Time Stamp       |  |
|----------|------|---|--------------------|----------------------|---------|------------------|--|
| <br>L1   | 5    | "251567".ap.  | US-PGPUB;<br>USPAT | AND                  | ON      | 2007/01/11 08:33 |  |
| L2       | 2    | "532273".ap.  | US-PGPUB;<br>USPAT | AND                  | ON      | 2007/01/11 09:45 |  |
| L3       | 6    | "821642".ap.  | US-PGPUB;<br>USPAT | AND                  | ON      | 2007/01/11 09:54 |  |
| L4       | 88   | 549/48.ccls.  | US-PGPUB;<br>USPAT | US-PGPUB; AND ON 200 |         |                  |  |
| L5       | 137  | 514/437.ccls.   | US-PGPUB;<br>USPAT | AND                  | ON ´    | 2007/01/11 09:55 |  |
| L6       | 6    | l4 and l5   | US-PGPUB;<br>USPAT | AND                  | ON      | 2007/01/11 09:56 |  |
| L7       | 173  | 549/461.ccls.   | US-PGPUB;<br>USPAT | AND                  | ON      | 2007/01/11 09:56 |  |
| L8       | 424  | 514/455.ccls.   | US-PGPUB;<br>USPAT | AND                  | ON      | 2007/01/11 09:56 |  |
| L9       | 1    | 17 and 18   | US-PGPUB;<br>USPAT | AND                  | ON      | 2007/01/11 09:56 |  |
| S1       | 0    | ("("6031122" "6359163" "6392078" <br>"RE37337").PN.").PN. | US-PGPUB;<br>USPAT | OR                   | OFF     | 2005/12/01 06:40 |  |
| S2       | . 4  | ("6031122" "6359163" "6392078" "<br>RE37337").PN.         | US-PGPUB;<br>USPAT | OR                   | ON      | 2005/12/01 06:49 |  |
| S3       | 0    | 588/277.icls  | US-PGPUB;<br>USPAT | OR .                 | ON      | 2005/12/01 06:50 |  |
| S4       | 0    | 588/277.ICLS  | US-PGPUB;<br>USPAT | OR                   | ON      | 2005/12/01 06:50 |  |
| S5       | 1    | 588/277.ICLS.   | US-PGPUB;<br>USPAT | OR                   | ON      | 2005/12/01 06:50 |  |
| S6       | 0    | 588/277ICLS.  | US-PGPUB;<br>USPAT | OR                   | ON      | 2005/12/01 06:50 |  |
| S7       | 1    | 588/277.ICLS.   | US-PGPUB;<br>USPAT | OR                   | ON      | 2005/12/01 06:51 |  |
| S8       | 0    | 588/277.cCLS.   | US-PGPUB;<br>USPAT | OR                   | ON      | 2005/12/01 06:51 |  |
| S9       | 0    | 588/277.CCLS.   | US-PGPUB;<br>USPAT | OR                   | ON      | 2005/12/01 06:52 |  |
| S10      | 274  | 558/277.CCLS.   | US-PGPUB;<br>USPAT | OR                   | ON      | 2005/12/01 06:53 |  |
| S11      | 208  | 558/277.iCLS.   | US-PGPUB;<br>USPAT | OR                   | ON      | 2005/12/01 06:55 |  |
| S12      | 0    | (RYU-yong OR Gelbein-Abraham).<br>IN.                     | US-PGPUB;<br>USPAT | OR                   | ON      | 2005/12/01 06:59 |  |

| S13 | 0              | RYU-yong.IN. OR Gelbein-Abraham.<br>IN. | US-PGPUB;<br>USPAT | OR    | ON   | 2005/12/01 06:59 |
|-----|----------------|---|--------------------|-------|------|------------------|
| S14 | 0              | RYU-yong.IN.                            | US-PGPUB;<br>USPAT | OR    | ON . | 2005/12/01 06:59 |
| S15 | 1565           | RYU.IN.                                 | US-PGPUB;<br>USPAT | OR    | ON   | 2005/12/01 07:01 |
| S16 | 0              | RYU-y.IN.                               | US-PGPUB;<br>USPAT | OR    | ON   | 2005/12/01 07:00 |
| S17 | 1565           | RYU.IN.                                 | US-PGPUB;<br>USPAT | OR    | ON   | 2005/12/01 07:00 |
| S18 | 0              | Gelbein-Abraham.IN.                     | US-PGPUB;<br>USPAT | OR    | ON   | 2005/12/01 07:00 |
| S19 | 55             | Gelbein.IN.                             | US-PGPUB;<br>USPAT | OR    | ON   | 2005/12/01 07:01 |
| S20 | <sub>.</sub> 4 | S17 AND S19                             | US-PGPUB;<br>USPAT | OR    | ON   | 2005/12/01 07:02 |
| S21 | 0              | S20 <"20030312"                         | US-PGPUB;<br>USPAT | AND   | ON   | 2005/12/01 07:05 |
| S22 | 2              | S20 AND @pd<="20030312"                 | US-PGPUB;<br>USPAT | AND   | ON   | 2005/12/01 07:14 |
| S23 | 0              | S20 AND @pd<="2003"                     | US-PGPUB;<br>USPAT | AND   | ON   | 2005/12/01 07:15 |
| S24 | 2              | S20 AND @pd<="20030312"                 | US-PGPUB;<br>USPAT | AND   | ON   | 2005/12/01 07:15 |
| S25 | 3              | S20 AND @py<="2003"                     | US-PGPUB;<br>USPAT | AND   | ON   | 2005/12/01 07:20 |
| S26 | 0              | "ryu-yong.in"                           | US-PGPUB;<br>USPAT | AND   | ON   | 2005/12/01 07:21 |
| S27 | 0              | "ryu.in"                                | US-PGPUB;<br>USPAT | AND   | ON   | 2005/12/01 07:22 |
| S28 | 1565           | ryu.IN.                                 | US-PGPUB;<br>USPAT | AND   | ON   | 2005/12/01 07:22 |
| S29 | 1565           | ryu.iN.                                 | US-PGPUB;<br>USPAT | AND   | ON   | 2005/12/01 07:22 |
| S30 | 1565           | ryu.in.                                 | US-PGPUB;<br>USPAT | AND   | ON   | 2005/12/01 07:22 |
| S31 | 0              | ryu-yong.in.                            | US-PGPUB;<br>USPAT | AND   | ON   | 2005/12/01 07:22 |
| S32 | 1565           | ryu.in.                                 | US-PGPUB;<br>USPAT | AND . | ON   | 2005/12/01 07:23 |
| S33 | 162            | yong.in. S32                            | US-PGPUB;<br>USPAT | AND   | ON   | 2005/12/01 07:23 |
| S34 | 25             | j.in. AND S33                           | US-PGPUB;<br>USPAT | AND   | ON   | 2005/12/01 07:24 |

| S35              | 18     | S34 AND @ad<="20030312"                           | US-PGPUB;<br>USPAT                          | AND  | ON   | 2005/12/01 07:26 |
|------------------|--------|---|---|------|------|------------------|
| S36              | 4      | ("6031122" "6359163" "6392078" "<br>RE37337").PN. | US-PGPUB;<br>USPAT                          | OR   | ON   | 2005/12/02 12:25 |
| S37              | 3      | ("4222944" "3759948" "5814651").<br>PN.           | US-PGPUB;<br>USPAT                          | OR   | ON   | 2005/12/02 13:14 |
| S38              | 0      | ("WO2004089940").PN.                              | USPAT;<br>USOCR                             | OR   | OFF  | 2005/12/02 13:17 |
| S39              |        | ("WO2004089940").PN.                              | US-PGPUB;<br>USPAT;<br>USOCR;<br>DERWENT    | OR   | OFF  | 2005/12/02 13:18 |
| 540              | 0      | ("104821642").PN.                                 | US-PGPUB;<br>USPAT                          | OR   | OFF  | 2005/12/04 07:17 |
| S41              | 0      | ("104821642.pn").PN.                              | US-PGPUB;<br>USPAT                          | OR   | OFF  | 2005/12/04 07:17 |
| S42              | .0     | ("10821642.pn").PN.                               | US-PGPUB;<br>USPAT                          | OR   | OFF  | 2005/12/04 07:17 |
| S43              | 1      | "10821642"  | US-PGPUB;<br>USPAT                          | OR   | ON.  | 2005/12/04 07:17 |
| S44              | 0      | WO037805  | US-PGPUB;<br>USPAT                          | OR   | ON   | 2005/12/04 12:35 |
| S45 <sup>'</sup> | 0      | WO2004/037805                                     | US-PGPUB;<br>USPAT                          | OR . | ON   | 2005/12/04 12:35 |
| S46              | 512015 | WO 2004/037805                                    | US-PGPUB;<br>USPAT                          | OR   | ON   | 2005/12/04 12:35 |
| S47              | 1      | 60/519,967  | US-PGPUB;<br>USPAT                          | AND  | ON   | 2005/12/05 15:20 |
| S48              | 0      | 459/461.ccls.                                     | US-PGPUB;<br>USPAT                          | AND  | ON   | 2005/12/06 08:03 |
| S49              | 172    | 549/461.ccls.                                     | US-PGPUB;<br>USPAT                          | AND  | ON   | 2005/12/06 08:04 |
| S50              | 100    | 549/461.ccls. AND<br>@ad<="20031113"              | US-PGPUB;<br>USPAT                          | AND  | ON   | 2005/12/06 08:06 |
| S51              | 50     | S50 AND dibenzofuran                              | US-PGPUB;<br>USPAT                          | AND  | ON . | 2005/12/06 08:11 |
| S52              | 0      | S51 AND "548".ccls.                               | US-PGPUB;<br>USPAT                          | AND  | ON   | 2005/12/06 08:12 |
| S53              | 5      | S51 AND 548/444.ccls.                             | US-PGPUB;<br>USPAT                          | AND  | ON.  | 2005/12/06 08:16 |
| S54              |        | S51 AND 548/444.ccls.                             | US-PGPUB;<br>USPAT;<br>EPO; JPO;<br>DERWENT | AND  | ON   | 2005/12/06 08:16 |

| S55 | 1   | "3846553".pn.                           | US-PGPUB;<br>USPAT                                    | OR    | ON   | 2006/05/11 11:14 |
|-----|-----|---|---|-------|------|------------------|
| S56 | 1   | "532273".ap.                            | US-PGPUB;<br>USPAT                                    | OR _  | ON   | 2006/05/11 11:16 |
| S57 | 0   | "532273".an.                            | US-PGPUB;<br>USPAT                                    | OR    | ON   | 2006/05/11 11:16 |
| S58 | 1   | "532273".ap.                            | US-PGPUB;<br>USPAT                                    | OR    | ON · | 2006/05/11 11:31 |
| S59 | 1   | "6110962".pn.                           | US-PGPUB;<br>USPAT                                    | OR    | ON   | 2006/05/11 11:31 |
| S60 | 3   | ("3759948" "3846553" "6110962").<br>PN. | US-PGPUB;<br>USPAT                                    | AND   | ON   | 2006/05/22 14:53 |
| S61 | 58  | 549/461.icls.                           | US-PGPUB;<br>USPAT;<br>USOCR;<br>EPO; JPO;<br>DERWENT | AND . | ON   | 2006/05/25 08:50 |
| S62 | 250 | 549/461.ccls.                           | US-PGPUB;<br>USPAT;<br>USOCR;<br>EPO; JPO;<br>DERWENT | AND . | ON   | 2006/05/25 08:50 |
| S63 | 266 | 546/284.7.ccls.                         | US-PGPUB;<br>USPAT;<br>USOCR;<br>EPO; JPO;<br>DERWENT | AND   | ON   | 2006/05/25 08:50 |
| S64 | 334 | 546/281.1.ccls.                         | US-PGPUB;<br>USPAT;<br>USOCR;<br>EPO; JPO;<br>DERWENT | AND   | ON   | 2006/05/25 08:51 |
| S65 | 289 | 549/43.ccls.                            | US-PGPUB;<br>USPAT;<br>USOCR;<br>EPO; JPO;<br>DERWENT | AND   | ON   | 2006/05/25 08:51 |
| S66 | 375 | 544/375.ccls.                           | US-PGPUB;<br>USPAT;<br>USOCR;<br>EPO; JPO;<br>DERWENT | AND   | ON   | 2006/05/25 08:52 |
| S67 | 333 | 544/153.ccls.                           | US-PGPUB;<br>USPAT;<br>USOCR;<br>EPO; JPO;<br>DERWENT | AND   | ON   | 2006/05/25 08:52 |

|      |    |                          | •   |     |      |                  |
|------|----|--------------------------|---|-----|------|------------------|
| ·S68 | 0  | S61 and S63              | US-PGPUB;<br>USPAT;<br>USOCR;<br>EPO; JPO;<br>DERWENT | AND | ON   | 2006/05/25 08:53 |
| S69  | 1  | S61 and S64              | US-PGPUB;<br>USPAT;<br>USOCR;<br>EPO; JPO;<br>DERWENT | AND | ON   | 2006/05/25 08:59 |
| S70  |    | S61 and S66              | US-PGPUB;<br>USPAT;<br>USOCR;<br>EPO; JPO;<br>DERWENT | AND | ON   | 2006/05/25 08:58 |
| S71  | 1. | S61 and S67              | US-PGPUB;<br>USPAT;<br>USOCR;<br>EPO; JPO;<br>DERWENT | AND | ON.  | 2006/05/25 08:53 |
| S72  |    | S62 and S63              | US-PGPUB;<br>USPAT;<br>USOCR;<br>EPO; JPO;<br>DERWENT | AND | ON   | 2006/05/25 08:57 |
| S73  | 1  | S62 and S64              | US-PGPUB;<br>USPAT;<br>USOCR;<br>EPO; JPO;<br>DERWENT | AND | ON   | 2006/05/25 08:59 |
| S74  | 14 | S62 and S66              | US-PGPUB;<br>USPAT;<br>USOCR;<br>EPO; JPO;<br>DERWENT | AND | ON   | 2006/05/25 08:59 |
| S75  | 21 | S62 and S67              | US-PGPUB;<br>USPAT;<br>USOCR;<br>EPO; JPO;<br>DERWENT | AND | ON . | 2006/05/25 09:02 |
| S76  | 0  | "03377"                  | JPO   | AND | ON   | 2006/08/15 06:04 |
| S77  | 0  | "03 <sup>3</sup> 377.ap" | JPO   | AND | ON   | 2006/08/15 06:05 |
| S78  | 0  | "3377"                   | JPO   | AND | ON   | 2006/08/15 06:05 |
| S79  | 0  | "014156"                 | JPO   | AND | ON   | 2006/08/15 06:05 |
| S80  | 1  | jp63014156               | JPO   | AND | ON   | 2006/08/15 06:06 |
| S81  | 1  | "63014156"               | JPO   | AND | ON . | 2006/08/15 06:06 |

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                 has been enhanced and reloaded
        OCT 30 CHEMLIST enhanced with new search and display field
NEWS 4
NEWS
        NOV 03
                 JAPIO enhanced with IPC 8 features and functionality
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        NOV 10
                 CA/CAplus F-Term thesaurus enhanced
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                 CAS Registry Number crossover limit increased to 300,000 in
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     8
                 additional databases
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        NOV 20
                 CA/CAplus to MARPAT accession number crossover limit increased
      9
                 to 50,000
NEWS 10
        DEC 01
                 CAS REGISTRY updated with new ambiguity codes
                 CAS REGISTRY chemical nomenclature enhanced
        DEC 11
NEWS 11
        DEC 14
                 WPIDS/WPINDEX/WPIX manual codes updated
NEWS 12
                 GBFULL and FRFULL enhanced with IPC 8 features and
NEWS 13
        DEC 14
                 functionality
        DEC 18
                 CA/CAplus pre-1967 chemical substance index entries enhanced
NEWS 14
                 with preparation role
NEWS 15
        DEC 18
                 CA/CAplus patent kind codes updated
NEWS 16
        DEC 18 MARPAT to CA/Caplus accession number crossover limit increased
                 to 50,000
NEWS 17
        DEC 18
                 MEDLINE updated in preparation for 2007 reload
        DEC 27
                 CA/CAplus enhanced with more pre-1907 records
NEWS 18
         JAN 08
                 CHEMLIST enhanced with New Zealand Inventory of Chemicals
NEWS 19
NEWS EXPRESS NOVEMBER 10 CURRENT WINDOWS VERSION IS V8.01c, CURRENT
              MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
              AND CURRENT DISCOVER FILE IS DATED 25 SEPTEMBER 2006.
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FILE 'HOME' ENTERED AT 07:49:35 ON 11 JAN 2007

=> file reg

COST IN U.S. DOLLARS . SINCE FILE TOTAL

ENTRY SESSION 0.21 0.21

FULL ESTIMATED COST

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http://www.cas.org/ONLINE/UG/regprops.html

Uploading C:\Documents and Settings\ychu\Desktop\Case\10821642\10821642b.str

chain nodes :

15 16 17 18 20

ring nodes :

1 2 3 4 5 6 7 8 9 10 11 12 13

chain bonds :

10-15 13-18 15-16 15-17

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-9 7-8 7-10 8-9 8-13 10-11 11-12 12-13

exact/norm bonds :

5-7 6-9 8-9 10-15 13-18 15-16 15-17

normalized bonds :

1-2 1-6 2-3 3-4 4-5 5-6 7-8 7-10 8-13 10-11 11-12 12-13

G1:0,S

### Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 13:Atom 15:CLASS 16:CLASS 17:CLASS 18:Atom 20:CLASS 21:Atom

#### L1 STRUCTURE UPLOADED

=> d

L1 HAS NO ANSWERS

L1 STR

Structure attributes must be viewed using STN Express query preparation.

=> s 11

SAMPLE SEARCH INITIATED 07:50:43 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 131 TO ITERATE

100.0% PROCESSED 131 ITERATIONS

4 ANSWERS

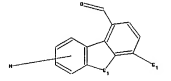
SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*

BATCH \*\*COMPLETE\*\*

PROJECTED ITERATIONS: 1934 TO 3306 PROJECTED ANSWERS: 4 TO 200 = >

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16 15

chain nodes :

15 16 17 19

ring nodes :

1 2 3 4 5 6 7 8 9 10 11 12 13

chain bonds :

10-15 13-17 15-16

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-9 7-8 7-10 8-9 8-13 10-11 11-12 12-13

exact/norm bonds :

5-7 6-9 8-9 10-15 13-17 15-16

normalized bonds :

1-2 1-6 2-3 3-4 4-5 5-6 7-8 7-10 8-13 10-11 11-12 12-13

G1:0,S

Match level :

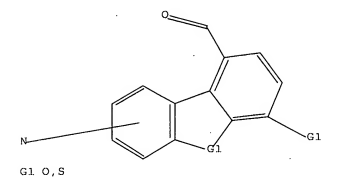
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 13:Atom 15:CLASS 16:CLASS 17:Atom 19:CLASS 20:Atom

L3 STRUCTURE UPLOADED

=> d

L3 HAS NO ANSWERS

L3 STR



Structure attributes must be viewed using STN Express query preparation.

=> s 13

SAMPLE SEARCH INITIATED 07:52:02 FILE 'REGISTRY'

SAMPLE SCREEN SEARCH COMPLETED - 417 TO ITERATE

100.0% PROCESSED 417 ITERATIONS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*

BATCH \*\*COMPLETE\*\* .

PROJECTED ITERATIONS: 7115 TO 9565

PROJECTED ANSWERS: 4 TO 200

L4 4 SEA SSS SAM L3

Uploading C:\Documents and Settings\ychu\Desktop\Case\10821642\10821642d.str

4 ANSWERS

chain nodes :

15 17

ring nodes :

1 2 3 4 5 6 7 8 9 10 11 12 13

chain bonds :

13-15

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-9 7-8 7-10 8-9 8-13 10-11 11-12 12-13

exact/norm bonds : 5-7 6-9 8-9 13-15

normalized bonds :

1-2 1-6 2-3 3-4 4-5 5-6 7-8 7-10 8-13 10-11 11-12 12-13

G1:0.S

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 13:Atom 15:Atom 17:CLASS 18:Atom

L5 STRUCTURE UPLOADED

=> d L5 HAS NO ANSWERS L5 STR

Structure attributes must be viewed using STN Express query preparation.

=> s 15

SAMPLE SEARCH INITIATED 07:52:44 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 7093 TO ITERATE

28.2% PROCESSED 2000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED) SEARCH TIME: 00.00.01 2 ANSWERS

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*

BATCH \*\*COMPLETE\*\*

PROJECTED ITERATIONS: 136811 TO 146909 PROJECTED ANSWERS: 2 TO 300

L6 2 SEA SSS SAM L5

=> s 15 full

FULL SEARCH INITIATED 07:52:59 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 142525 TO ITERATE

100.0% PROCESSED 142525 ITERATIONS

164 ANSWERS

SEARCH TIME: 00.00.03

L7 164 SEA SSS FUL L5

=> file caplus

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ENTRY SESSION

FULL ESTIMATED COST 173.90 174.11

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=> s 17

L8 23 L7

=> d ibib abs 15-23

L8 ANSWER 15 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1986:207666 CAPLUS Full-text

DOCUMENT NUMBER: 104:207666

TITLE: Peptide synthesis by prior thiol capture. 2. Design

of templates for intramolecular O,N-acyl transfer. 4,6-Disubstituted dibenzofurans as optimal spacing

elements

AUTHOR(S): Kemp, D. S.; Galakatos, Nicholas G.; Bowen, Benjamin;

Tan, Kenneth

CORPORATE SOURCE: Dep. Chem., Massachusetts Inst. Technol., Cambridge,

MA, 02139, USA

SOURCE: Journal of Organic Chemistry (1986), 51(10), 1829-38

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 104:207666

GI

AB A central feature of the strategy for amide bond formation by prior thiol capture is an intramol. acyl transfer across a template that links the phenolic ester function of one peptide with an unsym. disulfide involving the side chain of the N-terminal cysteine residue of a second peptide. The structures of 4-hydroxy-6-mercaptodibenzofuran (I) and 4-hydroxy-6-mercaptophenoxythiin (II) were established by 1H NMR spectra of deuterated dibenzofuran and phenoxythiin derivs. On the basis of the criterion of effective molarity, a dibenzofuran template for intramol. acyl transfer is approx. 2 orders of magnitude more efficient than a phenoxythiin. An effective local concn. of ca 5 M and a Hammett .rho. value of 2.6 is obsd. for the intramol. acyl-transfer reaction of O-acyl dibenzofuran derivs. III (R = Ac, PhCH2O2C-Ala; R1 = H, C1, Br, NO2; R2 = H, C1; R3 = C-terminal group) to the corresponding N-acyl derivs. IV.

L8 ANSWER 16 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1979:66597 CAPLUS Full-text

DOCUMENT NUMBER:

90:66597

TITLE:

Antiallergic activity of tetracyclic derivatives of

quinoline-2-carboxylic acid. 2. Some benzothienoquinolinecarboxylic acids

AUTHOR(S):

Wade, James J.; Erickson, Edward H.; Hegel, Ramon F.;

Lappi, Larry R.; Rice, Thomas K.

Riker Lab., 3M Co., St. Paul, MN, USA

SOURCE:

Journal of Medicinal Chemistry (1978), 21(9), 941-8

CODEN: JMCMAR; ISSN: 0022-2623

DOCUMENT TYPE:

CORPORATE SOURCE:

Journal

LANGUAGE:

English

GI

Benzothienoquinolinecarboxylic acids and their esters (>90 compds.) were AB prepd. and tested as potential antiallergic agents. Their antianaphylactic activity was comparable to that of di-Na cromoglycate. I and II were approx. 8 times more active than di-Na chromoglycate in rat passive cutaneous anaphylaxis assay.

ANSWER 17 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1977:423247 CAPLUS Full-text

DOCUMENT NUMBER:

87:23247

TITLE:

1-Benzothieno[3,2-f]quinolinecarboxylic acids

INVENTOR (S):

Lappi, Larry R.; Erickson, Edward H.

PATENT ASSIGNEE(S):

Riker Laboratories, Inc., USA

SOURCE:

Ger. Offen., 36 pp.

CODEN: GWXXBX

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO.             | KIND | DATE     | APPLICATION NO.   | DATE     |
|------------------------|------|----------|-------------------|----------|
|                        |      |          |                   |          |
| DE 2638081             | A1   | 19770310 | DE 1976-2638081   | 19760824 |
| US 4018780             | Α    | 19770419 | US 1975-607626    | 19750825 |
| JP 52027799            | Α    | 19770302 | JP 1976-101027    | 19760824 |
| FR 2321887             | A1   | 19770325 | FR 1976-25645     | 19760824 |
| FR 2321887             | B1   | 19781117 |                   |          |
| GB 1563112             | Α    | 19800319 | GB 1976-35227     | 19760824 |
| PRIORITY APPLN. INFO.: |      |          | US 197.5-607626 A | 19750825 |
| GI                     |      |          |                   |          |

1-Hydroxy[1]benzothieno[3,2-f]quinoline-3-carboxylic acids (I; R = e.g. 10-F, AB 8-Cl, 10-Br, 10-Me, 10-MeO, H, 8-MeO, 9-Me; R1 = e.g. H, 6-Cl, 5-MeO, 6-MeO; n= 0, 1, 2), useful as allergy inhibitors, are prepd. by reaction of 2aminodibenzothiophenes with MeO2CC.tplbond.CCO2Me (II), cyclization of the resulting di-Me (dibenzothiophene-2-ylamino) fumarates and hydrolysis of the Me esters. Thus, reaction of II with 2-amino-8-fluorodibenzothiophene in MeOH at room temp. 16 h gives di-Me [(8-fluorodibenzothiophene-2-yl)amino]fumarate which on heating 5 min at 240.degree. in Ph2O gives the Me ester which is hydrolyzed to give I (R = 10-F, R1 = H, n = 0).

ANSWER 18 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN ·L8 ACCESSION NUMBER:

DOCUMENT NUMBER:

1966:59731 CAPLUS Full-text 64:59731

ORIGINAL REFERENCE NO.: 64:11148g-h

TITLE:

Potentially chemotherapeutic dibenzofurans

AUTHOR(S): Onyiriuka, S. O.; Rees, A. H.

CORPORATE SOURCE: Univ. Ibadan, Nigeria

SOURCE: J. Chem. Soc., Org. (1966), (5), 504-6

DOCUMENT TYPE: Journal LANGUAGE: English

GI For diagram(s), see printed CA Issue.

AB A number of new disubstituted dibenzofurans (I and II) were prepd. for evaluation of their chemotherapy, and for use in further syntheses.

L8 ANSWER 19 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1966:59730 CAPLUS Full-text

DOCUMENT NUMBER: 64:59730
ORIGINAL REFERENCE NO.: 64:11148f-g

TITLE: The constituents of Cacalia decomposita. Structures of

maturin, maturinin, maturone, and maturinone

AUTHOR(S): Correa, J.; Romo, J.

CORPORATE SOURCE: Univ. Nacl. Autonoma, Mexico, D.F. SOURCE: Tetrahedron (1966), 22(2), 685-91

CODEN: TETRAB; ISSN: 0040-4020

DOCUMENT TYPE: Journal LANGUAGE: English

GI For diagram(s), see printed CA Issue.

The structures of maturin (I), maturnin (II), maturone (III), and maturinone (IV) have been established as furonaphthalene derivatives, closely related to cacalol (V) and cacalone (VI).

L8 ANSWER 20 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1962:457092 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 57:57092

ORIGINAL REFERENCE NO.: 57:11415i,11416a-b

TITLE: Neomycin-treated cellulosic textile materials

PATENT ASSIGNEE(S): American Cyanamid Co.

SOURCE: 8 pp.; Addn. to Brit. 788,968, (CA 52, 10601b)

SOURCE:

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

GB 900803 19620711 GB 1959-36877 19591030
PRIORITY APPLN. INFO.: US 19581103

AB A durable antibacterial finish can be applied to cellulosic textiles by treatment with neomycin B, neomycin C, and salts of the neomycin complex. The yellowness or dullness which usually appears upon laundring of neomycin (I)-treated fabricscan be counteracted by application of optical bleaching agents. Agents applicable to cellulosic textiles are acyldiaminostilbenes, triazinyldiaminostilbenes, and acyldiaminodibenzothiophene dioxides, all of which contain sulfonic acid groups. Other brighteners which can be used are the all-purpose types, such as benzimadazoles and triazoles. The treating soln. preferably a pad bath, contg. both I and the bleaching agent can be prepd. without copptn. of the ingredients by addn. of an alk. agent, such as NaOH or KOH, to the soln. The fabric is padded through the bath and dried at 150-350.degree.F. The concn. of I m the soln. can be varied within wide limits depending upon the intended use. When applied by padding, the soln. should contain 0.001-1% by wt. of I, and the same concn. of bleaching agent, preferably 0.01-0.05% of the dry wt. of the material.

L8 ANSWER 21 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1951:40188 CAPLUS Full-text

DOCUMENT NUMBER: 45:40188
ORIGINAL REFERENCE NO.: 45:6845c-g

TITLE: Relation between constitution and tinctorial

properties of substantive azoic dyes

AUTHOR(S): Krepelka, V.; Rais, J. CORPORATE SOURCE: Prague Polytech. Inst.

SOURCE: Collection of Czechoslovak Chemical Communications

(1950), 15, 412-32

CODEN: CCCCAK; ISSN: 0010-0765

DOCUMENT TYPE: Journal LANGUAGE: French

The substantivity (fs) and tinctorial power (v) to cotton of the following AB azoic dyestuffs have been detd. Main Component, .lambda.max.m.mu., emax., fs, v; aniline, 480, 38,275, 17.6, 0.64; 4,4'-diaminodiphenylamine, 525, 44,850, 36.55, 1.36; 4,4'-diaminodiphenylmethane, 497, 34,500, 22.45, 0.834; benzidine, 527.5, 40,350, 39.0, 1.42; 3,3'-dichlorobenzidine, 520, 47,000, 33.4, 1.33; benzidine-3,3'-disulfonic acid, 515, 94,000, 25.0, 1.17; benzidine-2,2'-disulfonic acid, 502.5, 78,600, 10.54, 0.474; benzidine sulfone, 540, 32,930, 19.9, 0.787; benzidine sulfone-3,3'-disulfonic acid, 540, 55,850, 14.31, 0.712; diamino-2,2'-stilbenedisulfonic acid, 532, 82,800, 42.1, 2.015; p-phenylenediamine (monoazo deriv.), 510, 15,610, 25.25, 0.96; pphenylenediamine (bisazo deriv.), 515, 43,300, 32.2, 1.05; p,p'diaminodiphenylurea, 491, 40,200, 33.15, 1.302; 2,2'-dinitro-4,4'diaminodiphenyl-methane, 497, 27,350, 12.2, 0.508; .omicron.-aminophenol, 500, 25,600, 20.6, 0.785; 3,3'-diamino-4,4'-dihydroxydiphenyl-methane, 495, 42,600, 26.75, 1.035; 3-aminosalicylic acid, 505, 34,500, 13.5, 0.604; 3,3'-diamino-5,5'-methylenedisalicylic acid, 502, 64,300, 18.55, 0.84; The dyestuffs were prepd. by coupling the diazotized main component with 6-amino-1-naphthol-3sulfonic acid. Substantivities were assigned numerical values and were detd. spectrophotometrically, titration with Ti salts, and colorimetrically. The following general rules were proposed for a bisazo dyestuff to be substantive: (1) the mol. wt. must be fairly high, (2) at least 2 auxochromes must be linked by a long chain of conjugated double bonds (at least 8), (3) free rotation of aromatic nuclei must be possible (thus dyestuffs from benzidine-2,2'-disulfonic acid are acid dyestuffs which dye wool), (4) usually the dyestuff should not be a deriv. of a p,p'-diamine, (5) neg. substituents decrease the substantivity.

L8 ANSWER 22 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1940:15386 CAPLUS Full-text

DOCUMENT NUMBER: 34:15386

ORIGINAL REFERENCE NO.: 34:2368g-i,2369a-i

TITLE: Dibenzofuran. XV. 1,4- and 1,4,6-Derivatives

AUTHOR(S): Gilman, Henry; Cheney, Lee C.

SOURCE: Journal of the American Chemical Society (1939), 61,

3149-56

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

Incidental to studies concerned with the bridging of the 1- and 9-positions in dibenzofuran types, series of 1,4- and 1,4,6-derivs. have been synthesized. In these compds. the 4- and 4,6-substituents are strong o,p-directors. 4- Hydroxy-6-methoxydibenzofuran (I) with HBr in AcOH gives 91.6% of 4,6- dihydroxydibenzofuran (II), m. 200-2.degree.. II (0.115 mol), 0.345 mol of Me2SO4, 39 mL. of 60% KOH and Me2CO give a nearly quant. yield of 4,6- dimethoxydibenzofuran (III), m. 128-9.degree.; picrate, deep yellow, m. 161-

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2.degree.. III and Accl with AlCl3 in PhNO2 give 60% of the 1-Ac deriv. (IV),
m. 178.5-9.5.degree.. The oxime of IV, m. 203-4.degree., with PCl5 in C6H6
gives 76.4% of 1-acetamino-4,6- dimethoxydibenzofuran, m. 244-5.degree.;
hydrolysis gives the HCl salt, m. 286-7.degree., of the 1-NH2 deriv. (V) of
III, m. 162-2.5.degree. I (4.28~g.) in 15~mL.~15% KOH and 385~mL.~H2O,
treated with the PhN2Cl from 0.02 mol PhNH2 at 3.degree., gives 56.6% of 1-
benzeneazo-4-hydroxy-6- methoxydibenzofuran, rust-colored, m. 175.degree.;
Me2SO4 and KOH in Me2CO give 88% of the 4,6-di-MeO deriv., deep orange, m.
170.degree.; redn. gives 32.8% of V. 4-Methoxydibenzofuran (VI) and 10%
excess of (COCl)2 in PhNO2, treated with 10% excess AlCl3 at 0.degree. and
allowed to stand at room temp. for 28 h., give a mixt. of 3 products; extn. of
the amorphous product with 5% NaOH gives 4-methoxy-1-dibenzofurancarboxylic
acid, m. 276-7.degree.; the alkali-insol. solid, extd. with AcOH, gives 18% of
bis(4-methoxy-1-dibenzofuryl) ketone, m. 234.degree.; the AcOH-insol. portion
is bi(4-methoxy-1-dibenzofuroyl), pale yellow, m. 329.degree., 34.6% yield.
VI and C1CH2COCl with AlCl3 in PhNO2 give 53.2% of the 1-chloroacetyl deriv.,
m. 165-6.degree.; ClCOCO2Et gives 43% of the 1-ethoxalyl deriv., m.
113.degree.; hydrolysis with 15% NaOH gives 4-methoxy-1-dibenzofuryl-.alpha.-
oxoacetic acid, pale yellow, m. 187.degree. (semicarbazone, m. 211.5-
12.degree. (decompn.)). III (4.56 g.) and (COCl)2 with AlCl3 in PhNO2 give
0.37 q. 4,6-dimechoxy-1- dibenzofurancarboxylic acid, m. 297-8.degree.; 10.4%
of bis(4,6-dimethoxy-1-benzofuryl) ketone, m. 254-5.degree., sol. in AcOH, and
60.7% of bi(4,6-dimethoxy-1-dibenzofuroyl), pale yellow, m. above 300.degree.,
insol. in AcOH. 3-Hydroxy-4-methoxydibenzofuran (VII) and HBr in AcOH give
88% of 3,4-dihydroxydibenzofuran (VIII), m. 164-4.5.degree.; di-Ac deriv., m.
104-5.degree.. VII and Me2SO4 with 10% NaOH give 81% of the 3,4-di-Me ether
(IX), m. 60-1.degree.. IX and AcCl give 55.5% of the 1-Ac deriv. (X), m.
90.5-1.degree.. The oxime of X, m. 156-7.degree., is rearranged by PCl5 in
C6H6 to give 94% of 1-acetamino-3,4-dimethoxydibenzofuran, m. 156-7.degree.;
hydrolysis gives 1-amino-3,4-dimethoxydibenzofuran, m. 162.5-3.degree., which
also results in 10% yield on heating the 1-Br deriv. of IX with concd. NH4OH
and CuBr for 14.5 h. at 220-30.degree.. III (22.8 g.) in 600 mL. AcOH and 100
mL. of M Br soln. in AcOH give 73% of the 1-Br deriv., m. 152.degree., and 12%
of a product m. 144-7.degree.; III (3 g.) and 52.7 mL. of a 0.5 M Br-AcOH
soln. give 74% of the 1,9-di-Br deriv. (XI), m. 167-8.degree.. II gives a
nearly quant. yield of the 1,9-di-Br deriv., m. 239-40.degree. (decompn.);
Me2SO4 gives XI. I gives 58.6% of the 1,3-di-Br deriv., m. 177-8.degree.;
Me2SO4 gives the 1,3-di-Br deriv. of III, m. 173.5-4.degree.. IX forms 88.5%
of the 1-Br deriv., m. 108.degree.; VII yields 54.6% of the 1-Br deriv., m.
161-2.degree., which was also prepd. from 1-bromo-3-amino-4-
methoxydibenzofuran through the diazo reaction in 21% yield. 4-Bromo-6-
methoxydibenzofuran (XII) and HI (d. 1.67) give 19% of the 6-HO analog, m.
138-9.degree.; FeCl3 gives a green color. XII, CuBr and NH4OH, heated in a
steel bomb for 10 h. at 100.degree. and for 8 h. at 215.degree., give 51% of
the HCl salt, m. 235-6.degree., of 4-amino-6-methoxydibenzofuran, m.
109.degree.; HBr in AcOH gives the 6-HO analog, m. 191.5-2.5.degree.. II,
NaHSO3 and concd. NH4OH, heated at 185-95.degree. for 20 h., give 81% of 4,6-
diaminodibenzofuran, m. 152.degree.; HCl salt, m. 298.degree. (decompn.);
picrate, red-brown, m. 213.degree. (decompn.); di-Ac deriv., m. 297-8.degree..
Di-Ac deriv. of II, m. 177.degree.. II and PhN2Cl give a dark brown compd.,
m. 228.degree. (decompn.), which is nearly pure 1,3,9-trisbenzeneazo deriv.;
Me2SO4 gives 77% of 1,3,9-trisbenzeneazo-4,6-dimethoxydibenzofuran, red-
orange, m. 191-3.degree.. The 1-Ac deriv. of III, oxidized with I-KI in NaOH-
dioxane, gives 55.2% of 4,6-dimethoxy-1-dibenzofurancarboxylic acid (XIII), m.
297-8.degree.; this also resulted from carbonation of the Grignard reagent of
the 1-Br deriv. of II; Me ester, m. 163.degree.. XIII gives an acid chloride,
m. 147-50.degree.; CH2N2 gives 21.2% of diazomethyl 4,6-dimethoxy-1-
dibenzofuryl ketone, pale yellow, m. 151.degree. (decompn.); heating the
ketone with concd. NH4OH and AgNO3 in dioxane gives 52% of the amide, m. 210-
11.degree., of 4,6-dimethoxy-1- dibenzofurylacetic acid, m. 205.5-6.5.degree..
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3-Aminodibenzofuran, diazotized and reduced with SnCl2, gives 87.3% of the HCl salt, m. 242-3.degree., of 3-hydrazinodibenzofuran, pale yellow, m. 174-5.degree., which turns orange in the atm. 4-Aminodibenzofuran in abs. EtOH, reduced by Na in a N atm., gives 62% of 1,2,3,4-tetrahydro-6-aminodibenzofuran, which is an oil at 0.degree.; HCl salt, pink, m. 228.degree. (decompn.); the diazo soln. with .beta.-Cl0H7OH gives a quant. yield of a brilliant carmine red dye, m. 199-201.degree..

L8 ANSWER 23 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1939:29848 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 33:29848
ORIGINAL REFERENCE NO.: 33:4238b-f

TITLE: Dibenzofuran. X. Aminohydroxy derivatives

AUTHOR(S): Gilman, Henry; Jacoby, Arthur L.; Swislowsky, Jack SOURCE: Journal of the American Chemical Society (1939), 61,

954-6

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

4-Acetaminodibenzofuran and HNO3 (d. 1.49) in Ac2O at -10.degree. give 35% of the Ac deriv. (I), pale yellow, m. 238.degree., of 3-nitro-4aminodibenzofuran (II), deep yellow, m. 185-6.degree.. Catalytic redn. of I gives a quant. yield of 3-amino-4-acetaminodibenzofuran, silvery plates, m. 236-7.degree.; di-Ac deriv., m. 257.degree.. Redn. of II with Raney Ni and reaction with phenanthraquinone give dibenzo[a,c]benzofuro[2,3- h]phenazine, yellow, m. 277-8.degree.. 4-Hydroxydibenzofuran (III) and HNO3 in AcOH at -12.degree. give 25% of the 3-NO2 deriv. (IV), light yellow, m. 193.degree.; this also results from II through the diazo reaction. IV and CH2N2 give 65% of the 4-MeO deriv., yellow, m. 129.5.degree.. Nitration of III with concd. HNO3 in AcOH at 60.degree. gives 77% of the 3,8-di-NO2 deriv. (V), orange-red, m. 225.degree. (decompn.); this also results in a nearly quant. yield from IV. V and CH2N2 give 83% of 3,8-dinitro-4-methoxydibenzofuran, orange, m. 177.degree.. The 2-isomer of III yields 80% of a yellow di-NO2 deriv., m. 240.degree. (decompn.), which is probably the 3,8-deriv. 4-Methoxydibenzofuran and HNO3 in Ac20 at -15.degree. to -20.degree. give 18% of the 1-NO2 deriv., m. 155.degree.; 1-NH2 deriv., pale lavender, m. 104.degree., which also results from 1-bromo-4-methoxydibenzofuran and concd. NH4OH with CuBr at 230-40.degree.. 4-Ethoxydibenzofuran gives 28% of the 1-NO2 deriv., yellow, m. 135-5.5.degree.; 1-NH2 deriv., m. 91.degree. (Ac deriv., m. 218.5.degree.). 3-Aminodibenzofuran, EtI, Na2CO3 and H2O, refluxed 48 h., give 70% of 3diethylaminodibenzofuran, m. 68.degree...

### => d ibib abs 1-14

L8 ANSWER 1 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2006:1283312 CAPLUS Full-text

DOCUMENT NUMBER: 146:35882

TITLE: Electrophotographic photoreceptor, process cartridge,

and electrophotographic apparatus

INVENTOR(S): Kikuchi, Toshihiro; Ochi, Atsushi; Sako, Harumi;

Yoshimura, Kimihiro; Tamai, Hideaki; Kosaka, Nobuo

PATENT ASSIGNEE(S): Canon Kabushiki Kaisha, Japan

SOURCE: PCT Int. Appl., 86pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

| PAT   | PATENT NO.  |      |     |     |     | KIND DATE |      |      | APPLICATION NO. |                  |      |      |     | DATE     |      |      |     |
|-------|-------------|------|-----|-----|-----|-----------|------|------|-----------------|------------------|------|------|-----|----------|------|------|-----|
|       |             |      |     |     |     | -         |      |      |                 |                  |      |      |     |          |      |      |     |
| WO    | 2006        | 1298 | 79  |     | A1  |           | 2006 | 1207 | 1               | WO 2006-JP311464 |      |      |     | 20060601 |      |      |     |
|       | W:          | ΑE,  | AG, | AL, | AM, | AT,       | ΑÜ,  | AZ,  | BA,             | BB,              | ·BG, | BR,  | BW, | BY,      | ΒZ,  | CA,  | CH, |
|       |             | CN,  | CO, | CR, | CŪ, | CZ,       | DE,  | DK,  | DM,             | DZ,              | EC,  | EE,  | EG, | ES,      | FI,  | GB,  | GD, |
|       |             | GE,  | GH, | GM, | HR, | HU,       | ID,  | IL,  | IN,             | IS,              | ΚE,  | KG,  | KM, | KN,      | ΚP,  | KR,  | KZ, |
|       |             | LC,  | LK, | LR, | LS, | LT,       | LU,  | LV,  | LY,             | MA,              | MD,  | MG,  | MK, | MN,      | MW,  | MX,  | MZ, |
|       | NA, NG, NI, |      | NO, | NZ, | OM, | PG,       | PH,  | PL,  | PT,             | RO,              | RU,  | SC,  | SD, | SE,      | SG,  |      |     |
|       |             | SK,  | SL, | SM, | SY, | TJ,       | TM,  | TN,  | TR,             | TT,              | TZ,  | UA,  | ÜĠ, | US,      | UZ,  | VC,  | VN, |
|       |             | YU,  | ZA, | ZM, | ZW  |           |      |      |                 |                  |      |      |     |          |      |      | •   |
|       | RW:         | AT,  | BE, | BG, | CH, | CY,       | CZ,  | DE,  | DK,             | EE,              | ES,  | FI,  | FR, | GB,      | GR,  | HU,  | IE, |
|       |             | IS,  | IT, | LT, | LU, | LV,       | MC,  | NL,  | PL,             | PT,              | RO,  | SE,  | SI, | SK,      | TR,  | BF,  | ВJ, |
|       |             | CF,  | CG, | CI, | CM, | GA,       | GN,  | GQ,  | GW,             | ML,              | MR,  | NE,  | SN, | TD,      | TG,  | BW,  | GH, |
|       |             | GM,  | KE, | LS, | MW, | MZ,       | NA,  | SD,  | SL,             | SZ,              | TZ,  | UG,  | ZM, | ZW,      | AM,  | AZ,  | BY, |
|       |             | KG,  | KZ, | MD, | RU, | TJ,       | TM   |      |                 |                  | •    |      |     |          |      |      |     |
| ORTTY | KG, KZ, MD, |      |     |     |     |           |      |      |                 | TP 2             | 005- | 1627 | 3.0 |          | A 20 | 0050 | 602 |

PRIORITY APPLN. INFO.:

JP 2005-162730 A 20050602 JP 2005-162732 A 20050602

This invention provides an electrophotog. photoreceptor, which, while ensuring satisfactory mech. strength, has significantly improved charge transport properties and has satisfactory elec. characteristics, and a process cartridge and an electrophotog. app. In the electrophotog. photoreceptor, the outermost surface layer comprises at least a product produced by polymg. or crosslinking a charge transport triarylamine compd. contg. a chain polymerizable functional group and curing the polymn. or crosslinking product. The process cartridge and electrophotog. app. comprise the photoreceptor.

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 2 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2006:361248 CAPLUS Full-text

2

DOCUMENT NUMBER:

144:412351

TITLE:

Process for the preparation of n-(3,5-dichloropyridin-

4-yl)-4-difluoromethoxy-8-methanesulfonamido-

dibenzo[b,d]furan-1-carboxamide

INVENTOR(S):

Gopalan, Balasubramanian; Gharat, Laxmikant Atmaram; Chandrasekhar, Batchu; Karaunakaran, Usha; Pillai,

Bijukumar Gopinathan

PATENT ASSIGNEE(S):

Glenmark Pharmaceuticals S.A., Switz.

SOURCE:

PCT Int. Appl., 54 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT N      | PATENT NO. |        |             |             | DATE |     |     | APPLICATION NO. |     |     |     |     | DATE     |     |     |
|---------------|------------|--------|-------------|-------------|------|-----|-----|-----------------|-----|-----|-----|-----|----------|-----|-----|
|               |            |        |             | -           |      |     |     |                 |     |     |     |     |          |     |     |
| WO 20060      | 40652      |        | A2          | A2 20060420 |      |     | 1   | WO 2005-IB3035  |     |     |     |     | 20051012 |     |     |
| WO 2006040652 |            |        | A3 20061026 |             |      |     |     |                 |     |     |     |     |          |     |     |
| W:            | AE, AG     | G, AL, | AM,         | AT,         | AU,  | AZ, | BA, | BB,             | BG, | BR, | BW, | BY, | ΒZ,      | CA, | CH, |
|               | CN, CC     | O, CR, | CU,         | CZ,         | DE,  | DK, | DM, | DZ,             | EC, | EE, | EG, | ES, | FI,      | GB, | GD, |
|               | GE, GH     | H, GM, | HR,         | HU,         | ID,  | IL, | IN, | IS,             | JP, | KE, | KG, | KM, | KΡ,      | KR, | KZ, |
|               | LC, LK     | K, LR, | LS,         | LT,         | LU,  | LV, | LY, | MA,             | MD, | MG, | MK, | MN, | MW,      | MX, | MZ, |
|               | NA, NO     | G, NI, | NO,         | NZ,         | OM,  | PG, | PH, | PL,             | PT, | RO, | RU, | SC, | SD,      | SE, | SG, |
|               | SK, SI     | L, SM, | SY,         | TJ,         | TM,  | TN, | TR, | TT,             | TZ, | UA, | UG, | US, | UZ,      | VC, | VN, |
|               | YU, ZA     | A, ZM, | zw          |             |      |     |     |                 |     |     |     |     |          | •   |     |
| RW:           | AT, BE     | E, BG, | CH,         | CY,         | CZ,  | DE, | DK, | EE,             | ES, | FI, | FR, | GB, | GR,      | HU, | IE, |

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IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ,
             CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH,
             GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
             KG, KZ, MD, RU, TJ, TM
                                20060622
                                            US 2005-251567
     US 2006135779
                         A1
                                                                   20051013
PRIORITY APPLN. INFO.:
                                            US 2004-618474P
                                                                P 20041013
                                            IN 2004-MU1099
                                                                A 20041014
                                            US 2004-621981P
                                                                P 20041021
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The present invention relates to a method of prepg. N-(3,5-dichloropyridin- 4-yl)-4-difluoromethoxy-8-methanesulfonamido-dibenzo[b,d] furan-1- carboxamide and pharmaceutically acceptable salts thereof, such as its sodium salt, and novel intermediate compds. useful in the synthesis of the aforementioned compd. For example, reaction of 4-cyclopentyloxy-3- hydroxybenzaldehyde with 2-bromo-1-fluoro-4-nitrobenzene (70-77%), followed by cyclization, gave 4-cyclopentyloxy-8-nitro-1- formyldibenzofuran in 60-65% yield, which yielded the title compd. after 9 steps.

L8 ANSWER 3 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2006:103883 CAPLUS Full-text

DOCUMENT NUMBER:

144:170874

TITLE:

Preparation of dibenzofurans and related compounds as phosphodiesterase type 4 inhibitors useful for the treatment of inflammatory and allergic disorders

INVENTOR (S):

Balasubramanian, Gopalan; Gharat, Laxmikant Atmaram;

Joshi, Hemant Vasant

PATENT ASSIGNEE(S):

Glenmark Pharmaceuticals Ltd., India

SOURCE:

PCT Int. Appl., 145 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

Engr

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT      | NO.       |         |         |                 | APPL    | CATION 1  | 10.     | DATE        |  |  |  |
|-------------|-----------|---------|---------|-----------------|---------|-----------|---------|-------------|--|--|--|
|             |           |         |         |                 |         |           |         |             |  |  |  |
| WO 20.0     | 6011024   | A2      | 2006    | 0202            | WO 20   | 005-IB206 | 51.     | 20050718    |  |  |  |
| WO 200      | 6011024   | A3      | 2006    | 0330            |         | •         |         |             |  |  |  |
| W:          | AE, AG,   | AL, AM, | AT, AU, | AZ, I           | BA, BB, | BG, BR,   | BW, BY, | BZ, CA, CH, |  |  |  |
|             | CN, CO,   | CR, CU, | CZ, DE, | DK, I           | DM, DZ, | EC, EE,   | EG, ES, | FI, GB, GD, |  |  |  |
|             | GE, GH,   | GM, HR, | HU, ID, | IL,             | IN, IS, | JP, KE,   | KG, KM, | KP, KR, KZ, |  |  |  |
|             | LC, LK,   | LR, LS, | LT, LU, | LV, I           | MA, MD, | MG, MK,   | MN, MW, | MX, MZ, NA, |  |  |  |
|             | NG, NI,   | NO, NZ, | OM, PG, | PH,             | PL, PT, | RO, RU,   | SC, SD, | SE, SG, SK, |  |  |  |
|             | SL, SM,   | SY, TJ, | TM, TN, | TR,             | TT, TZ, | UA, UG,   | US, UZ, | VC, VN, YU, |  |  |  |
|             | ZA, ZM,   | ZW      |         |                 |         |           |         |             |  |  |  |
| RW          | : AT, BE, | BG, CH, | CY, CZ, | DE, I           | DK, EE, | ES, FI,   | FR, GB, | GR, HU, IE, |  |  |  |
|             | IS, IT,   | LT, LU, | LV, MC, | NL,             | PL, PT, | RO, SE,   | SI, SK, | TR, BF, BJ, |  |  |  |
|             | CF, CG,   | CI, CM, | GA, GN, | GQ, (           | GW, ML, | MR, NE,   | SN, TD, | TG, BW, GH, |  |  |  |
|             | GM, KE,   | LS, MW, | MZ, NA, | SD, S           | SL, SZ, | TZ, UG,   | ZM, ZW, | AM, AZ, BY, |  |  |  |
|             | KG, KZ,   | MD, RU, | TJ, TM  |                 |         |           |         |             |  |  |  |
| PRIORITY AP | PLN. INFO | .:      |         | US 2004-589479P |         |           | 79P     | P 20040719  |  |  |  |
| ,           |           |         |         |                 | IN 20   | 004-MU809 | €       | A 20040729  |  |  |  |

OTHER SOURCE(S): MARPAT 144:170874

GΙ

The present invention relates to novel tricyclic compds. (shown as I; AB variables defined below; e.g. 4-(4-methoxydibenzofuran-1-yl)-2pyrrolidinone), analogs, tautomers, regioisomers, stereoisomers, enantiomers, diastereomers, polymorphs, pharmaceutically acceptable salts, appropriate oxides, pharmaceutically acceptable solvates and pharmaceutical compns. contg. them. The present invention also relates to phosphodiesterase type 4 (PDE4) inhibitors which down regulate or inhibit the prodn. of TNF-.alpha. and therefore are useful in the treatment of variety of allergic and inflammatory diseases including asthma and chronic obstructive pulmonary disease (COPD). Methods of prepn. are claimed and prepns. and/or characterization data for .apprx.60 examples of I are included. For example, 4-(4-methoxydibenzofuran-1-y1)-2-pyrrolidinone was prepd. by reductive cyclization of 3-(4methoxydibenzofuran-1-yl)-4- nitrobutanoate (prepn. given) in iPrOH/DMF using 10 % Pd/C. For I: R1 is (un) substituted aryl, arylalkyl, heteroaryl, heterocyclyl, heterocyclylalkyl, or heteroarylalkyl; R2, R3, R4, R5 and R6 may be the same or different and = H or (un) substituted alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkylalkyl, cycloalkenyl, aryl, arylalkyl, heteroaryl, heterocyclic group, heterocyclylalkyl, or heteroarylalkyl, -NR8R9, -C(O)R8, -C(O)OR8, -C(O)NR8R9, -S(O)mR8, -S(O)mNR8R9, nitro, -OH, cyano, formyl, acetyl, halogen, -OR8, -SR8, or a protecting group, or when R1 and R3, or R4 and R5 are ortho to each other then R1 and R3 together with the C atoms to which they are bound or R4 and R5 together with the C atoms to which they are bound may be joined to a form a (un)satd. cyclic ring, which may optionally include up to two heteroatoms = O, NRa or S; X is O, S(O)m and NR6; P is O or S; m = 0-2; addnl. details are given in the claims. IC50 values for inhibition of PDE4 by .apprx.60 examples of I are tabulated.

ANSWER 4 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER:

2005:1077670 CAPLUS Full-text

DOCUMENT NUMBER:

143:372827

TITLE:

Hair dyeing compositions containing heterocyclic

INVENTOR(S):

Glenn, Robert Wayne; McMeekin, Anthony; Lim, Mu'ill; Gardlik, John Michael; Jones, Stevan David; Murphy,

Bryan Patrick

PATENT ASSIGNEE(S):

UK

SOURCE:

U.S. Pat. Appl. Publ., 42 pp.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO.     | KIND   | DATE         | APPLICATION NO.       | DATE        |  |
|----------------|--------|--------------|-----------------------|-------------|--|
|                |        |              |                       |             |  |
| US 2005217038  | A1     | 20051006     | US 2005-99740         | 20050406    |  |
| WO 2005099656  | A2     | 20051027     | 20050406              |             |  |
| WO 2005099656  | A3     | 20060504     | •                     |             |  |
| W: AE, AG, AL, | AM, AT | , AU, AZ, BA | , BB, BG, BR, BW, BY, | BZ, CA, CH, |  |

CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG EP 1735059 20061227 EP 2005-737700 A2 R: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LI, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR PRIORITY APPLN. INFO.: US 2004-559823P P 20040406 WO 2005-US11810 . W 20050406

OTHER SOURCE(S): MARPAT 143:372827

AB Compns. for the oxidative dyeing of hair fibers, comprise a medium suitable for dyeing and 1 or more tricyclic heteroarom. hair dyeing compds. A method for oxidative dyeing of the hair fibers, comprises applying such compns. in the presence of an oxidizing agent, for a period sufficient to develop the desired coloration. Thus, a hair dye formulation contained 9H-carbazole-1,4-diamine 0.05%.

L8 ANSWER 5 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2005:1026087 CAPLUS Full-text

DOCUMENT NUMBER:

143:336429

TITLE:

The color conversion film, the color conversion filter

and the organic EL display

INVENTOR(S):

Yoshida, Shohei; Oyama, Yosuke; Kawaguchi, Takeshi;

Kobayashi, Makoto

PATENT ASSIGNEE(S):

Kochi University, Japan; Fuji Electric Holding Co.,

Ltd.

SOURCE:

Jpn. Kokai Tokkyo Koho, 25 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent .

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO.             | KIND   | DATE       | AP | PLICATION NO. |   | DATE     |
|------------------------|--------|------------|----|---------------|---|----------|
|                        |        |            |    |               |   |          |
| JP 2005259688          | Α      | 20050922   | JP | 2005-34904    |   | 20050210 |
| PRIORITY APPLN. INFO.: |        |            | JP | 2004-37033    | Α | 20040213 |
| OTHER SOURCE(S):       | MARPAT | 143:336429 |    |               |   |          |

GI

### \* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

The disclosed color conversion film contains a binder and a dye selected from I or II (R1 = alkyl, cycloalkyl, phenyl; R2, R3 = C1-10 alkyl, C5-10 cycloalkyl; R4, R5 = H; R2-R4 and R3-R5 combination may form alkylene groups to complete rings; X = alkyl, cycloalkyl, Ph, halo, OR6, R6CO2, SR6, NR6R7; R6, R7 = H, alkyl cycloalkyl; Z = O, S, NR6). Optionally, the film may also contain red-conversion dye selected from rhodamine, cyanine, pyridine, and oxazine dyes. Color filters and org. electroluminescent display devices which

uses the color conversion films are also disclosed. The film exhibit excellent light-fastness.

L8 ANSWER 6 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:99226 CAPLUS Full-text

DOCUMENT NUMBER: 142:197859

TITLE: Preparation of dibenzo[b,f]furan-1-carboxamides,

9H-carbazole-4-carboxamides, and dibenzo[b,d]thiophene-4-carboxamides as PDE4 inhibitors for the treatment of

inflammatory and allergic disorders

INVENTOR(S): Gopalan, Balasubramanian; Gharat, Laxmikant A.;

Lakdawala, Aftab D.; Karunakaran, Usha

PATENT ASSIGNEE(S): Glenmark Pharmaceuticals, Inc. USA, USA

SOURCE: U.S. Pat. Appl. Publ., 59 pp., Cont.-in-part of Appl.

No. PCT/IB04/000355.

CODEN: USXXCO

DOCUMENT TYPE:

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

| PAT      | PATENT NO.            |       |     |     | KIND DATE |          | APPLICATION NO. |               |                 |      |      | DATE |            |            |      |     |     |    |
|----------|-----------------------|-------|-----|-----|-----------|----------|-----------------|---------------|-----------------|------|------|------|------------|------------|------|-----|-----|----|
|          |                       |       |     |     |           | -        |                 |               |                 |      |      |      |            |            | -    |     |     |    |
| US :     | 2005                  | 0271  | 29  |     | A1        | 20050203 |                 |               | US 2004-821642  |      |      |      | 20040409   |            |      |     |     |    |
| WO :     | 2004                  | 08994 | 40  |     | A1        | :        | 2004            | 1021          | WO 2004-IB355   |      |      |      | 20040211   |            |      |     |     |    |
|          | <b>W</b> :            | ΑE,   | AG, | AL, | AM,       | AT,      | AU,             | AZ,           | BA,             | BB,  | BG,  | BR,  | BW,        | BY,        | ΒZ,  | CA, | CH, |    |
|          |                       | CN,   | CO, | CR, | CU,       | CZ,      | DE,             | DK,           | DM,             | DZ,  | EC,  | EE,  | EG,        | ES,        | FI,  | GB, | GD, |    |
|          |                       | GE,   | GH, | GM, | HR,       | HU,      | ID,             | IL,           | IN,             | IS,  | JP,  | KE,  | KG,        | ΚP,        | KR,  | ΚZ, | LC, |    |
|          |                       | LK,   | LR, | LS, | LT,       | LU,      | LV,             | MA,           | MD,             | MG,  | MK,  | MN,  | MW,        | MX,        | MZ,  | NA, | NI, |    |
|          |                       | NO,   | NZ, | OM, | PG,       | PH,      | PL,             | PT,           | RO,             | RU,  | SC,  | SD,  | SE,        | SG,        | SK,  | SL, | SY, |    |
|          |                       | TJ,   | TM, | TN, | TR,       | TT,      | TZ,             | ÜΑ,           | ΰĠ,             | US,  | UZ,  | VC,  | VN,        | YU,        | ZA,  | ZM, | zw  |    |
|          | RW:                   | BW,   | GH, | GM, | KE,       | LS,      | MW,             | MZ,           | SD,             | SL,  | SZ,  | TZ,  | UG,        | ZM,        | ZW,  | AM, | AZ, |    |
|          |                       | BY,   | KG, | ΚŻ, | MD,       | RU,      | TJ,             | TM,           | ΑT,             | BE,  | BG,  | CH,  | CY,        | CZ,        | DE,  | DK, | EE, |    |
|          |                       | ES,   | FI, | FR, | GB,       | GR,      | HU,             | ΙE,           | IT,             | LU,  | MC,  | NL,  | PT,        | RO,        | SE,  | SI, | SK, |    |
|          |                       | TR,   | BF, | ВJ, | CF,       | CG,      | CI,             | CM,           | GA,             | GN,  | GQ,  | GW,  | ML,        | MR,        | NE,  | SN, | TD, | TG |
| PRIORITY | RIORITY APPLN. INFO.: |       |     | . : |           |          |                 | IN 2003-MU363 |                 |      |      | 7    | A 20030411 |            |      |     |     |    |
|          |                       |       |     |     |           |          |                 |               | US 2003-519967P |      |      |      | ,          | P 20031113 |      |     |     |    |
|          |                       |       |     |     |           |          |                 | 1             | WO 2            | 004- | IB35 | 5    | 1          | A2 2       | 0040 | 211 |     |    |

OTHER SOURCE(S): MARPAT 142:197859

GI

Title heterocyclic tricycles I [wherein R1-R3, R5, R6, Ra = independently H, AB (un) substituted (cyclo) alkyl, (cyclo) alkenyl, alkynyl, (hetero) aryl, heterocyclyl(alkyl), etc.; R4 = NR5R6 (R5, R6 = H, alkyl, cycloalkyl, etc.), heterocyclyl; Ar = (un) substituted aryl(alkyl), heterocyclyl, heteroaryl; X = O, SO0-2, NRa; Y = CONR7, NR7SO0-2, SO0-2NR7, NR7CO; R7 = H, OH, ORa, (un) substituted alkyl, aryl, heterocyclyl; P = O, S; m = 0-3; n = 1-4; Ra = H, alkyl, cycloalkyl, etc.; and tautomers, regioisomers, stereoisomers, enantiomers, diastereomers, polymorphs, N-oxides, pharmaceutically acceptable salts, solvates, and compns. thereof] were prepd. as phosphodiesterase type 4 (PDE4) inhibitors. For example, N-(3,5-dichloropyrid-4-yl)-4-methoxy-8aminodibenzo[b,f]furan-1- carboxamide (prepd. in six steps from isovanillin, 4-fluoronitrobenzene, and 4-amino-3,5-dichloropyridine) was coupled with methanesulfonyl chloride in THF and pyridine to give the sulfonamide II. latter inhibited the PDE4-induced conversion of [3H] cAMP to the corresponding [3H] 5'-AMP with IC50 of 0.5058 nM. Thus, I and their pharmaceutical compns. are useful for the treatment of immune disorders, inflammatory conditions, allergic conditions, CNS diseases, and insulin resistant diabetes (no data).

L8 ANSWER 7 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2004:878393 CAPLUS Full-text

DOCUMENT NUMBER:

141:366121

TITLE:

Preparation of dibenzo[b, f] furan-1-carboxamides,

9H-carbazole-4-carboxamides, and dibenzo[b,d]thiophene-4-carboxamides as PDE4 inhibitors for the treatment of

inflammatory and allergic disorders

INVENTOR(S):

Gopalan, Balasubramanian; Gharat, Laxmikant Atmaram;

Lakdawala, Aftab Dawoodbhai; Karaunakaran, Usha

PATENT ASSIGNEE(S): Glenmark Pharmaceuticals Ltd., India

SOURCE:

PCT Int. Appl., 121 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

Engil

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

WO 2004089940 A1 20041021 WO 2004-IB355 20040211

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,

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CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
             GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC,
             LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI,
             NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY,
             TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
         RW: BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ,
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             TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
     AU 2004228453
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                                 20041021
                                             AU 2004-228453
                                                                     20040211
     CA 2522023
                          A1
                                 20041021
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             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
                                 20060509
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                                                                     20040211
     BR 2004009747
                          Α
                                             CN 2004-80016048
                                                                     20040211
     CN 1829711
                          Α
                                 20060906
     JP 2006522789
                          Т
                                 20061005
                                             JP 2006-506259
                                                                     20040211
     US 2005027129
                          A1
                                 20050203
                                             US 2004-821642
                                                                     20040409
                                                                     20051110
                                 20060111
                                             NO 2005-5316
     NO 2005005316
                          A
                                             IN 2003-MU363
                                                                     20030411
PRIORITY APPLN. INFO.:
                                             US 2003-519967P
                                                                     20031113
                                             WO 2004-IB355
                                                                  W .
                                                                     20040211
                         CASREACT 141:366121; MARPAT 141:366121
OTHER SOURCE(S):
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$$(R^3)$$
 m  $(R^4)$  n  $(R^4$ 

GΙ

Title heterocyclic tricycles I [wherein R1-R3, R5, R6, Ra = independently H, (un) substituted (cyclo) alkyl, (cyclo) alkenyl, alkynyl, (hetero) aryl, heterocyclyl (alkyl), etc.; R4 = NR5R6, heterocyclyl; Ar = (un) substituted aryl (alkyl), heterocyclyl, heteroaryl; X = O, S00-2, NRa; Y = CONR7, NR7SO0-2, S00-2NR7, NR7CO; R7 = H, OH, ORa, (un) substituted alkyl, aryl, heterocyclyl; P = O, S; m = 0-3; n = 1-4; and tautomers, regioisomers, stereoisomers, enantiomers, diastereomers, polymorphs, N-oxides, pharmaceutically acceptable salts, solvates, and compns. thereof] were prepd. as phosphodiesterase type 4 (PDE4) inhibitors. For example, N-(3,5-dichloropyrid-4-yl)-4-methoxy-8-aminodibenzo[b,f] furan-1- carboxamide (prepd. in six steps from isovanillin, 4-fluoronitrobenzene, and 4-amino-3,5-dichloropyridine) was coupled with methanesulfonyl chloride in THF and pyridine to give the sulfonamide II. The

latter inhibited the PDE4-induced conversion of [3H] cAMP to the corresponding [3H] 5'-AMP with IC50 of 0.5058 nM. Thus, I and their pharmaceutical compns. are useful for the treatment of immune disorders, inflammatory conditions, allergic conditions, CNS diseases, and insulin resistant diabetes (no data).

REFERENCE COUNT:

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 8 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2004:606600 CAPLUS Full-text

DOCUMENT NUMBER:

141:164539

TITLE:

Backlight polar organic light-emitting device

INVENTOR(S): PATENT ASSIGNEE(S): Lazarev, Pavel I. Optiva, Inc., USA

SOURCE:

PCT Int. Appl., 45 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

|      | PATENT NO.    |      |      | KIND DATE |     | APPLICATION NO. |     |      |      |     | DATE |       |                |     |     |     |         |     |
|------|---------------|------|------|-----------|-----|-----------------|-----|------|------|-----|------|-------|----------------|-----|-----|-----|---------|-----|
|      |               |      |      |           | -   |                 |     |      |      |     |      |       | <del>-</del> - | -   |     |     |         |     |
|      | WO            | 2004 | 0641 | 12        | •   | A2              |     | 2004 | 0729 |     | WO 2 | 004-1 | US22           | 9   |     | 2   | 0040    | 106 |
|      | WO 2004064112 |      |      | A3        |     | 20050317        |     |      | •    |     |      |       |                |     |     |     |         |     |
|      |               | W:   | AE,  | AG,       | AL, | AM,             | AT  | AU,  | AZ,  | BA, | BB,  | BG,   | BR,            | ВW, | BY, | ΒZ, | CA,     | CH, |
|      |               |      | CN,  | CO,       | CR, | CU,             | CZ, | DE,  | DK,  | DM, | DZ,  | EC,   | EE,            | EG, | ES, | FI, | GB,     | GD, |
|      |               |      | GE,  | GH,       | GM, | HR,             | HU, | ID,  | IL,  | IN, | IS,  | JP,   | KE,            | KG, | ΚP, | KR, | ΚZ,     | LC, |
|      |               |      | LK,  | LR,       | LS, | LT,             | LU, | LV,  | MA,  | MD, | MG,  | MK,   | MN,            | MW, | MX, | ΜZ  |         |     |
|      | US            | 2004 | 2241 | 82        |     | Al              |     | 2004 | 1111 |     | US 2 | 003-  | 6432           | 57  |     | 2   | 0 0,3 0 | 818 |
|      | JΡ            | 2006 | 5168 | 14        |     | T               |     | 2006 | 0706 |     | JP 2 | 006-  | 5008           | 03  |     | 2   | 0040    | 106 |
| PRIO | RITY          | APP  | LN.  | INFO      | . : |                 |     |      |      |     | US 2 | 003-  | 4387           | 14P |     | P 2 | 0030    | 107 |
|      |               |      |      |           |     |                 |     |      |      |     | US 2 | 003-  | 6432           | 57  |     | A 2 | 0030    | 818 |
|      |               |      |      |           |     |                 |     |      |      |     | WO 2 | 004-1 | US22           | 9   | 1   | W 2 | 0040    | 106 |

An org. light-emitting device (OLED) is provided which comprises a substrate and an org. electroluminescent cell formed on the substrate. The org. electroluminescent cell comprises a first electrode that serves as an anode, a second electrode that serves as a cathode, and at least one light-emitting layer positioned between the anode and cathode. At least one light-emitting layer is an anisotropic elec. conducting layer which has a globally ordered cryst. structure and is comprised of rodlike supramols.

ANSWER 9 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2004:370918 CAPLUS Full-text

DOCUMENT NUMBER:

140:391192

TITLE:

Preparation of dibenzofuran/dibenzothiophene

derivatives useful for the treatment of inflammatory

and allergic disorders

INVENTOR(S):

Balasubramanian, Gopalan; Gharat, Laxmikant Atmaram;

Lakdawala, Aftab Dawoodbhai; Anupindi, Raghu Ram

PATENT ASSIGNEE(S):

Glenmark Pharmaceuticals Ltd., India

PCT Int. Appl., 254 pp. SOURCE: CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

```
20040506
                                            WO 2003-IB4442
                                                                    20031008
    WO 2004037805
                          A1
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE,
             GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK,
             LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ,
             OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM,
             TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
             KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE. ES.
             FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
             BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                                            CA 2003-2503015
    CA 2503015
                          A1
                                20040506
                                                                    20031008
                          Al
                                20040513
                                            AU 2003-269317
                                                                    20031008
    AU 2003269317
                                20050720
                                            EP 2003-751096
    EP 1554262
                          Al
                                                                    20031008
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
                                20050802
                                           BR 2003-14721
     BR 2003014721
                          Α
                                                                    20031008
                                20060201
                                            CN 2003-80107246
                                                                    20031008
    CN 1729181
                          Α
                          Т
                                20060223
                                            JP 2004-546246
                                                                    20031008
    JP 2006506379
     ZA 2005002969
                          Α
                                20060222
                                            ZA 2005-2969
                                                                    20050413
     US 2006178418
                          A1
                                20060810
                                            US 2005-532273
                                                                    20050926
PRIORITY APPLN. INFO.:
                                            IN 2002-MU922
                                                                A 20021023
                                                               W 20031008
                                            WO 2003-IB4442
```

OTHER SOURCE(S):

MARPAT 140:391192

GI

Title compds. I [R1-3 = H, alk(en/yn)yl, cycloalkyl, etc.; P = 0, S; n = 0-4; AR Ar = (un)substituted aryl, etc.; Y = carboxamido, aminosulfonyl, etc.] are prepd. For instance, 4-methoxydibenzofuran-1-carboxylic acid (prepn. given) is converted to the corresponding acid chloride (PhH, SOCl2, reflux, 4 h) and treated with 4-amino-3,5-dichloropyridine (DMF/THF, NaH, -10.degree.) to give II. II has IC50 = 0.8 nM for PDE4. I are useful for the treatment of inflammatory conditions, diseases of the central nervous and insulin resistant diabetes.

CAPLUS COPYRIGHT 2007 ACS on STN ANSWER 10 OF 23

ACCESSION NUMBER: 2002:861050 CAPLUS Full-text

DOCUMENT NUMBER: 139:164660

Product class 6: dibenzothiophenes TITLE:

Andrews, M. D. AUTHOR(S):

Pfizer Central Research, Kent, CT13 9NJ, UK CORPORATE SOURCE: Science of Synthesis (2001), 10, 211-263 SOURCE:

CODEN: SSCYJ9

PUBLISHER: Georg T DOCUMENT TYPE: Journal

Georg Thieme Verlag
Journal; General Review

LANGUAGE:

English

AB A review. Methods for prepg. dibenzothiophenes are reviewed including

cyclization, ring transformation, aromatization and substituent modifications.

REFERENCE COUNT:

THERE ARE 187 CITED REFERENCES AVAILABLE FOR

THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L8 ANSWER 11 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN

187

ACCESSION NUMBER:

2000:694445 CAPLUS Full-text

DOCUMENT NUMBER:

133:268224

TITLE:

Methine dyes which are effectively excited with helium-neon laser and give strong fluorescence

INVENTOR(S):

Nishigaki, Junji; Kobayashi, Masaru; Kato, Takashi

PATENT ASSIGNEE(S):

Fuji Photo Film Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 19 pp.

SOURCE:

Jpn. Kokai Tokkyo Koho, 19 pr CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

GΙ

Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO.             | KIND   | DATE         | APPLICATION NO. | DATE     |  |
|------------------------|--------|--------------|-----------------|----------|--|
|                        |        |              |                 |          |  |
| JP 2000273330          | A      | 20001003     | JP 1999-79493   | 19990324 |  |
| PRIORITY APPLN. INFO.: |        |              | JP 1999-79493   | 19990324 |  |
| OTHER SOURCE(S):       | MARPAT | r·133:268224 |                 |          |  |

AB Title dyes are represented by the general formula I, where R = (un)substituted alkyl or (un)substituted aryl; Q = methine or polymethine having 1 or .gtoreq.2 substituents selected from (un)substituted hetero ring and (un)substituted aryl; and V11-16 = H or monovalent substituent. Substances labeled with I are also claimed.

L8 ANSWER 12 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2000:253229 CAPLUS Full-text

DOCUMENT NUMBER:

133:73821

TITLE:

Synthesis, structure and properties of

2-bromo-4,4'-dinitro-3'-(dimethylsulfonio)-2'-

biphenolate

AUTHOR(S):

Hou, Zi-Jie; Cai, Li-Ping

CORPORATE SOURCE:

National Laboratory of Applied Organic Chemistry,
Institute of Organic Chemistry, Lanzhou University,

Lanzhou, 730000, Peop. Rep. China

SOURCE:

Huaxue Xuebao (2000), 58(3), 358-362

CODEN: HHHPA4; ISSN: 0567-7351

PUBLISHER: Kexue Chubanshe

DOCUMENT TYPE: Journal LANGUAGE: Chinese

OTHER SOURCE(S): CASREACT 133:73821

In presence of an amine, the reaction of 3,7-dinitrodibenzobromolium bisulfate with DMSO gives 2-bromo-4,4'-dinitro-3'-dimethylsulfinio-2'- biphenolate (I). Crystal structure of I.1/2 C6H6, obtained by crystn. of I in benzene showes that I is a inner salt. The dihedral angle between two Ph rings is 117.3.degree.. The C(3)-O(3) bond distance(0.1254 nm) falls in the range from 0.119 .apprx. 0.121 nm (C=O double bond distance) to 0.133 .apprx. 0.136 nm (C-O single bond distance), which showes that there is a conjugation between 0(3) and the Ph ring. Thermal decompn. of I gives both 2-bromo-2'-methoxy-3'methylthio-4,4'-dinitro-biphenyl and 3,7-dinitro-4-methylthio-dibenzofurane. The reaction of I with hydrochloric acid gives corresponding sulfonium salt.

ANSWER 13 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1992:485229 CAPLUS Full-text

DOCUMENT NUMBER: 117:85229

TITLE: Triazine derivatives as plant virucides

Hofferek, Horst; Noll, Bernd; Keil, Siegfried; INVENTOR(S):

Kochmann, Werner; Ostermann, Wolf Dieter Chemie A.-G. Bitterfeld-Wolfen, Germany PATENT ASSIGNEE(S):

Ger. (East), 7 pp. SOURCE:

CODEN: GEXXA8

DOCUMENT TYPE: Patent

LANGUAGE: German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO.             | KIND   | DATE      | APPLICATION NO. | DAŤE     |  |
|------------------------|--------|-----------|-----------------|----------|--|
|                        | ~      |           |                 |          |  |
| DD 298196              | A5     | 19920213  | DD 1989-326011  | 19890224 |  |
| PRIORITY APPLN. INFO.: |        |           | DD 1989-326011  | 19890224 |  |
| OTHER SOURCE(S):       | MARPAT | 117:85229 |                 |          |  |
| GI                     |        |           |                 |          |  |

AB The triazines I [R1, R3 = C1, OH, (un) substituted NH2, heterocyclyl, etc.; R2, R4 = R1, Q, Q1, etc.; X = Na, K, NH4; Z = H, C1, SO3X; R5, R6 = H, SO3Na; R6R6 = SO2] are plant virucides, which enhance plant resistance against viruses. I

(R1 = R3 = NHCH2CH2OH, R2 = R4 = 3-NaO3SC6H4, R5 = H, R6 = SO3Na) (0.05%)protected tobacco against artificial infection by the tobacco mosaic virus.

ANSWER 14 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1990:138901 CAPLUS Full-text

DOCUMENT NUMBER:

112:138901

TITLE:

Preparation of nitrosodibenzofuran derivatives as dye

intermediates

INVENTOR(S):

Yamamoto, Shinichi; Taniguchi, Takashi

PATENT ASSIGNEE(S): SOURCE:

Toray Industries, Inc., Japan Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO.             | KIND   | DATE       | APPLICATION NO. | DATE     |
|------------------------|--------|------------|-----------------|----------|
|                        |        |            |                 |          |
| JP 01193261            | Α      | 19890803   | JP 1988-18138   | 19880128 |
| PRIORITY APPLN. INFO.: | ,      |            | JP 1988-18138   | 19880128 |
| OTHER SOURCE(S):       | MARPAT | 112:138901 | *               |          |

GI

$$R^{2}m$$
 $R^{3}n$ 

AR The title compds. [I; R1 = (substituted) NH2, OH, C2-10 acyloxy, C1-20 alkoxy, etc.; R2, R3 = (substituted) NH2, OH, C2-10 acyloxy, halo, cyano, etc.; m = 0-4; n = 0-2], useful as intermediates for dyes, are prepd. by nitrosation of dibenzofuran derivs. A 20% aq. NaNO2 soln. was added to a soln. of 10 g 2hydroxydibenzofuran in pyridine at 0.degree., followed by 30% H2SO4 with stirring, to give 11 g nitroso deriv. I (R1 = OH; m = n = 0). Similarly prepd. were 3 addnl. I.

#### => d ibib abs hitstr 15-23

ANSWER 15 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1986:207666 CAPLUS Full-text

DOCUMENT NUMBER:

104:207666

TITLE:

Peptide synthesis by prior thiol capture. 2. Design of templates for intramolecular O,N-acyl transfer. 4,6-Disubstituted dibenzofurans as optimal spacing

elements

AUTHOR (S):

Kemp, D. S.; Galakatos, Nicholas G.; Bowen, Benjamin;

Tan, Kenneth

CORPORATE SOURCE:

Dep. Chem., Massachusetts Inst. Technol., Cambridge,

MA, 02139, USA

SOURCE:

Journal of Organic Chemistry (1986), 51(10), 1829-38

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 104:207666

A central feature of the strategy for amide bond formation by prior thiol AΒ capture is an intramol. acyl transfer across a template that links the phenolic ester function of one peptide with an unsym. disulfide involving the side chain of the N-terminal cysteine residue of a second peptide. The structures of 4-hydroxy-6-mercaptodibenzofuran (I) and 4-hydroxy-6mercaptophenoxythiin (II) were established by 1H NMR spectra of deuterated dibenzofuran and phenoxythiin derivs. On the basis of the criterion of effective molarity, a dibenzofuran template for intramol. acyl transfer is approx. 2 orders of magnitude more efficient than a phenoxythiin. An effective local concn. of ca 5 M and a Hammett .rho. value of 2.6 is obsd. for the intramol. acyl-transfer reaction of O-acyl dibenzofuran derivs. III (R = Ac, PhCH2O2C-Ala; R1 = H, Cl, Br, NO2; R2 = H, Cl; R3 = C-terminal group) to the corresponding N-acyl derivs. IV.

101762-25-8P IT

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn. and O-acylation of, with alanine deriv.)

RN 101762-25-8 CAPLUS

Carbono(dithioperoxoic) acid, SS-(6-hydroxy-9-nitro-4-dibenzofuranyl) CN O-methyl ester (9CI) (CA INDEX NAME)

IT 101762-26-9P 101774-02-1P

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of)

RN 101762-26-9 CAPLUS

CN L-Alanine, N-[(phenylmethoxy)carbonyl]-, 6-[(methoxycarbonyl)dithio]-1-nitro-4-dibenzofuranyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 101774-02-1 CAPLUS

CN L-Alanine, N-[(1,1-dimethylethoxy)carbonyl]-3-[(6-hydroxy-9-nitro-4-dibenzofuranyl)dithio]-, methyl ester, ester with N[(phenylmethoxy)carbonyl]-L-alanine (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L8 ANSWER 16 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1979:66597 CAPLUS Full-text

DOCUMENT NUMBER:

90:66597

TITLE:

Antiallergic activity of tetracyclic derivatives of

quinoline-2-carboxylic acid. 2. Some benzothienoquinolinecarboxylic acids

AUTHOR(S):

Wade, James J.; Erickson, Edward H.; Hegel, Ramon F.;

Lappi, Larry R.; Rice, Thomas K.

CORPORATE SOURCE:

Riker Lab., 3M Co., St. Paul, MN, USA

SOURCE:

Journal of Medicinal Chemistry (1978), 21(9), 941-8

CODEN: JMCMAR; ISSN: 0022-2623

DOCUMENT TYPE:

Journal

LANGUAGE:

English

GI

Benzothienoquinolinecarboxylic acids and their esters (>90 compds.) were AB prepd. and tested as potential antiallergic agents. Their antianaphylactic activity was comparable to that of di-Na cromoglycate. I and II were approx. 8 times more active than di-Na chromoglycate in rat passive cutaneous anaphylaxis assay.

IT 67086-01-5P

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn. and cyclization of)

RN 67086-01-5 CAPLUS

2-Butenedioic acid, 2-[(6-methoxy-2-dibenzothienyl)amino]-, dimethyl ester CN(CA INDEX NAME)

IT 62986-33-8P

> RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of)

62986-33-8 CAPLUS RN

2-Dibenzothiophenamine, 6-methoxy- (9CI) (CA INDEX NAME) CN

Elvest priar

CAPLUS COPYRIGHT 2007 ACS on STN ANSWER 17 OF 23

ACCESSION. NUMBER:

1977:423247 CAPLUS Full-text

DOCUMENT NUMBER:

INVENTOR(S):

87:23247

TITLE:

1-Benzothieno[3,2-f]quinolinecarboxylic acids

Lappi, Larry R.; Erickson, Edward H.

PATENT ASSIGNEE(S):

Riker Laboratories, Inc., USA

SOURCE:

Ger. Offen., 36 pp.

DOCUMENT TYPE:

CODEN: GWXXBX Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

#### PATENT INFORMATION:

| PATENT NO.             | KIND | DATE ·   | APPLICATION NO.  | DATE     |
|------------------------|------|----------|------------------|----------|
|                        |      |          |                  |          |
| DE 2638081             | Al · | 19770310 | DE 1976-2638081  | 19760824 |
| US 4018780             | A    | 19770419 | US_ 1975-607626. | 19750825 |
| JP 52027799            | A    | 19770302 | JP 1976-101027   | 19760824 |
| FR 2321887             | A1   | 19770325 | FR 1976-25645    | 19760824 |
| FR 2321887             | Bl   | 19781117 |                  |          |
| GB 1563112             | A    | 19800319 | GB 1976-35227    | 19760824 |
| PRIORITY APPLN. INFO.: | 0    |          | US 1975-607626 A | 19750825 |

- 1-Hydroxy[1]benzothieno[3,2-f]quinoline-3-carboxylic acids (I; R = e.g. 10-F, 8-Cl, 10-Br, 10-Me, 10-MeO, H, 8-MeO, 9-Me; Rl = e.g. H, 6-Cl, 5-MeO, 6-MeO; n = 0, 1, 2), useful as allergy inhibitors, are prepd. by reaction of 2-aminodibenzothiophenes with MeO2CC.tplbond.CCO2Me (II), cyclization of the resulting di-Me (dibenzothiophene-2-ylamino)fumarates and hydrolysis of the Me esters. Thus, reaction of II with 2-amino-8-fluorodibenzothiophene in MeOH at room temp. 16 h gives di-Me [(8-fluorodibenzothiophene-2-yl)amino]fumarate which on heating 5 min at 240.degree. in Ph2O gives the Me ester which is hydrolyzed to give I (R = 10-F, Rl = H, n = 0).
  - 62986-47-4P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn. and cyclization of)

RN 62986-47-4 CAPLUS

IT

CN 2-Butenedioic acid, 2-[(6-methoxy-2-dibenzothienyl)amino]-, dimethyl ester, (Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

- IT 62986-33-8P
  - RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of, and reaction with dimethyl acetylenedicarboxylate)

RN 62986-33-8 CAPLUS

CN 2-Dibenzothiophenamine, 6-methoxy- (9CI) (CA INDEX NAME)

ANSWER 18 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN L8 1966:59731 CAPLUS Full-text ACCESSION NUMBER: DOCUMENT NUMBER: 64:59731 ORIGINAL REFERENCE NO.: 64:11148g-h Potentially chemotherapeutic dibenzofurans AUTHOR (S): Onyiriuka, S. O.; Rees, A. H. Univ. Ibadan, Nigeria CORPORATE SOURCE: J. Chem. Soc., Org. (1966), (5), 504-6 SOURCE: DOCUMENT TYPE: Journal English LANGUAGE: For diagram(s), see printed CA Issue. A number of new disubstituted dibenzofurans (I and II) were prepd. for AΒ evaluation of their chemotherapy, and for use in further syntheses. 5917-25-9P, 4-Dibenzofuranol, 8-nitro- 5917-26-0P, IT 4-Dibenzofuranol, 7-nitro- 5917-27-1P, Dibenzofuran, 6-methoxy-2-nitro- 5917-28-2P, Dibenzofuran, 6-methoxy-3-nitro-5918-98-9P, Dibenzofuran, 1-bromo-4-methoxy-8-nitro-5919-00-6P, 2-Dibenzofuranamine, 6-methoxy- 5919-01-7P, 3-Dibenzofuranamine, 6-methoxy- 5919-02-8P, Dibenzofuran, 2-acetamido-6-methoxy- 5919-03-9P, Dibenzofuran, 3-acetamido-6-methoxy- 5919-09-5P, Ketone, 4-methoxy-7-nitro-1dibenzofuranyl methyl 5919-10-8P, Ketone, 4-methoxy-8-nitro-1dibenzofuranyl methyl 5919-11-9P, Ketone, 4-methoxy-8-nitro-1dibenzofuranyl methyl, oxime 5919-12-0P, 1-Dibenzofurancarboxamide, 4-methoxy-N-methyl-7-nitro- 5946-69-0P, 1-Dibenzofurancarboxylic acid, 4-methoxy-7-nitro- 5981-98-6P, 4-Dibenzofuranol, 8-acetamido-, acetate 5981-99-7P, 4-Dibenzofuranol, 7-acetamido-, acetate 5982-01-4P, Ketone, 4-methoxy-7-nitro-1-dibenzofuranyl methyl, oxime 6077-63-0P, Ketone, 7-amino-4-methoxy-1-dibenzofuranyl methyl RL: PREP (Preparation)

RN

CN

(prepn. of) 5917-25-9 CAPLUS

RN 5917-26-0 CAPLUS
CN 4-Dibenzofuranol, 7-nitro- (7CI, 8CI) (CA INDEX NAME)

4-Dibenzofuranol, 8-nitro- (7CI, 8CI) (CA INDEX NAME)

RN 5917-27-1 CAPLUS

CN Dibenzofuran, 6-methoxy-2-nitro- (7CI, 8CI) (CA INDEX NAME)

RN 5917-28-2 CAPLUS

CN Dibenzofuran, 6-methoxy-3-nitro- (7CI, 8CI) (CA INDEX NAME)

RN 5918-98-9 CAPLUS

CN Dibenzofuran, 1-bromo-4-methoxy-8-nitro- (7CI, 8CI) (CA INDEX NAME)

RN 5919-00-6 CAPLUS

CN 2-Dibenzofuranamine, 6-methoxy- (7CI, 8CI) (CA INDEX NAME)

RN 5919-01-7 CAPLUS

CN 3-Dibenzofuranamine, 6-methoxy- (7CI, 8CI) (CA INDEX NAME)

RN 5919-02-8 CAPLUS

CN Dibenzofuran, 2-acetamido-6-methoxy- (7CI, 8CI) (CA INDEX NAME)

RN 5919-03-9 CAPLUS

CN Dibenzofuran, 3-acetamido-6-methoxy- (7CI, 8CI) (CA INDEX NAME)

RN 5919-09-5 CAPLUS

CN Ketone, 4-methoxy-7-nitro-1-dibenzofuranyl methyl (7CI, 8CI) (CA INDEX NAME)

RN 5919-10-8 CAPLUS

CN Ketone, 4-methoxy-8-nitro-1-dibenzofuranyl methyl (7CI, 8CI) (CA INDEX NAME)

RN 5919-11-9 CAPLUS

CN Ketone, 4-methoxy-8-nitro-1-dibenzofuranyl methyl, oxime (7CI, 8CI) (CA INDEX NAME)

RN 5919-12-0 CAPLUS

CN 1-Dibenzofurancarboxamide, 4-methoxy-N-methyl-7-nitro- (7CI, 8CI) (CA INDEX NAME)

RN 5946-69-0 CAPLUS

CN 1-Dibenzofurancarboxylic acid, 4-methoxy-7-nitro- (7CI, 8CI) (CA INDEX NAME)

RN 5981-98-6 CAPLUS

CN 4-Dibenzofuranol, 8-acetamido-, acetate (7CI, 8CI) (CA INDEX NAME)

5981-99-7 CAPLUS RN

4-Dibenzofuranol, 7-acetamido-, acetate (7CI, 8CI) (CA INDEX NAME) CN

RN 5982-01-4 CAPLUS

Ketone, 4-methoxy-7-nitro-1-dibenzofuranyl methyl, oxime (7CI, 8CI) (CA CNINDEX NAME)

6077-63-0 CAPLUS RN

Ketone, 7-amino-4-methoxy-1-dibenzofuranyl methyl (7CI, 8CI) (CA INDEX CN

ANSWER 19 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN L8

ACCESSION NUMBER:

1966:59730 CAPLUS Full-text

DOCUMENT NUMBER:

64:59730

ORIGINAL REFERENCE NO.: 64:11148f-g

TITLE:

The constituents of Cacalia decomposita. Structures of

maturin, maturinin, maturone, and maturinone

AUTHOR(S):

Correa, J.; Romo, J.

CORPORATE SOURCE:

Univ. Nacl. Autonoma, Mexico, D.F. Tetrahedron (1966), 22(2), 685-91

CODEN: TETRAB; ISSN: 0040-4020

DOCUMENT TYPE:

Journal

LANGUAGE:

SOURCE:

English

For diagram(s), see printed CA Issue. GI

The structures of maturin (I), maturnin (II), maturone (III), and maturinone AB (IV) have been established as furonaphthalene derivatives, closely related to cacalol (V) and cacalone (VI).

5917-25-9P, 4-Dibenzofuranol, 8-nitro- 5917-26-0P, IT

4-Dibenzofuranol, 7-nitro- 5919-00-6P, 2-Dibenzofuranamine,

6-methoxy- 5919-01-7P, 3-Dibenzofuranamine, 6-methoxy-

5919-09-5P, Ketone, 4-methoxy-7-nitro-1-dibenzofuranyl methyl

5919-10-8P, Ketone, 4-methoxy-8-nitro-1-dibenzofuranyl methyl

5919-11-9P, Ketone, 4-methoxy-8-nitro-1-dibenzofuranyl methyl,

oxime 5919-12-0P, 1-Dibenzofurancarboxamide,

4-methoxy-N-methyl-7-nitro- 5946-69-0P, 1-Dibenzofurancarboxylic

acid, 4-methoxy-7-nitro- 5981-98-6P, 4-Dibenzofuranol,

8-acetamido-, acetate 5981-99-7P, 4-Dibenzofuranol,

7-acetamido-, acetate 5982-01-4P, Ketone, 4-methoxy-7-nitro-1-

dibenzofuranyl methyl, oxime 6077-63-0P, Ketone,

7-amino-4-methoxy-1-dibenzofuranyl methyl

RL: PREP (Preparation)

(prepn. of)

RN 5917-25-9 CAPLUS

CN 4-Dibenzofuranol, 8-nitro- (7CI, 8CI) (CA INDEX NAME)

RN 5917-26-0 CAPLUS

4-Dibenzofuranol, 7-nitro- (7CI, 8CI) (CA INDEX NAME) CN

5919-00-6 CAPLUS RN

2-Dibenzofuranamine, 6-methoxy- (7CI, 8CI) (CA INDEX NAME) CN

RN 5919-01-7 CAPLUS

CN 3-Dibenzofuranamine, 6-methoxy- (7CI, 8CI) (CA INDEX NAME)

RN 5919-09-5 CAPLUS

CN Ketone, 4-methoxy-7-nitro-1-dibenzofuranyl methyl (7CI, 8CI) (CA INDEX NAME)

RN 5919-10-8 CAPLUS

CN Ketone, 4-methoxy-8-nitro-1-dibenzofuranyl methyl (7CI, 8CI) (CA INDEX NAME)

RN 5919-11-9 CAPLUS

CN Ketone, 4-methoxy-8-nitro-1-dibenzofuranyl methyl, oxime (7CI, 8CI) (CA INDEX NAME)

RN 5919-12-0 CAPLUS

CN 1-Dibenzofurancarboxamide, 4-methoxy-N-methyl-7-nitro- (7CI, 8CI) (CA INDEX NAME)

RN 5946-69-0 CAPLUS

CN 1-Dibenzofurancarboxylic acid, 4-methoxy-7-nitro- (7CI, 8CI) (CA INDEX NAME)

RN 5981-98-6 CAPLUS

CN 4-Dibenzofuranol, 8-acetamido-, acetate (7CI, 8CI) (CA INDEX NAME)

RN 5981-99-7 CAPLUS

CN 4-Dibenzofuranol, 7-acetamido-, acetate (7CI, 8CI) (CA INDEX NAME)

RN 5982-01-4 CAPLUS

CN Ketone, 4-methoxy-7-nitro-1-dibenzofuranyl methyl, oxime (7CI, 8CI) (CA INDEX NAME)

RN 6077-63-0 CAPLUS

CNKetone, 7-amino-4-methoxy-1-dibenzofuranyl methyl (7CI, 8CI) (CA INDEX

ANSWER 20 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN L8

ACCESSION NUMBER:

1962:457092 CAPLUS Full-text

DOCUMENT NUMBER:

57:57092

ORIGINAL REFERENCE NO.: 57:11415i,11416a-b

DATE

TITLE:

Neomycin-treated cellulosic textile materials

PATENT ASSIGNEE(S):

American Cyanamid Co.

SOURCE:

8 pp.; Addn. to Brit. 788,968, (CA 52, 10601b)

APPLICATION NO.

DATE

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

KIND

PATENT INFORMATION:

PATENT NO.

| GB 900803                  | 19620711        | GB 1959-368   | 77 19591030                   |
|----------------------------|-----------------|---------------|-------------------------------|
| PRIORITY APPLN. INFO.:     |                 | US            | 19581103                      |
| AB A durable antibacteria  | l finish can be | applied to    | cellulosic textiles by        |
| treatment with neomycin    | n B, neomycin C | , and salts   | of the neomycin complex. The  |
| yellowness or dullness     | which usually   | appears upon  | laundring of neomycin (I)-    |
| treated fabricscan be      | counteracted by | application   | of optical bleaching agents.  |
| Agents applicable to co    | ellulosic texti | les are acyl  | diaminostilbenes,             |
| triazinyldiaminostilbe     | nes, and acyldi | aminodibenzo  | chiophene dioxides, all of    |
| which contain sulfonic     | acid groups.    | Other bright  | eners which can be used are   |
| the all-purpose types,     | such as benzim  | adazoles and  | triazoles. The treating       |
| soln. preferably a pad     | bath, contg. b  | oth I and the | e bleaching agent can be      |
| prepd. without copptn.     | of the ingredi  | ents by addn  | . of an alk. agent, such as   |
| NaOH or KOH, to the so     | ln. The fabric  | is padded t   | nrough the bath, and dried at |
| 150-350.degree.F. The      | concn. of I m   | the soln. car | n be varied within wide       |
| limits depending upon      | the intended us | e. When app   | lied by padding, the soln.    |
| should contain 0.001-19    | by wt. of I,    | and the same  | concn. of bleaching agent,    |
| preferably 0.01-0.05% (    | of the dry wt.  | of the mater  | ial.                          |
| IT 103006-30-0P, 4,6-Diben | zothiophenedisu | ulfonic acid, |                               |

3,7-di-p-anisamido-, 5,5-dioxide

RL: PREP (Preparation)

(bleaching of neomycin-treated textiles by)

103006-30-0 CAPLUS RN

4,6-Dibenzothiophenedisulfonic acid, 3,7-di-p-anisamido-, 5,5-dioxide CN (CA INDEX NAME)

CAPLUS COPYRIGHT 2007 ACS on STN ANSWER 21 OF 23

1951:40188 CAPLUS Full-text ACCESSION NUMBER:

45:40188 DOCUMENT NUMBER:

ORIGINAL REFERENCE NO.: 45:6845c-g

Relation between constitution and tinctorial TITLE:

properties of substantive azoic dyes

Krepelka, V.; Rais, J. AUTHOR (S):

CORPORATE SOURCE: Prague Polytech. Inst.

Collection of Czechoslovak Chemical Communications SOURCE:

(1950), 15, 412-32

CODEN: CCCCAK; ISSN: 0010-0765

Journal DOCUMENT TYPE:

French LANGUAGE: The substantivity (fs) and tinctorial power (v) to cotton of the following AB azoic dyestuffs have been detd. Main Component, .lambda.max.m.mu., emax., fs, v; aniline, 480, 38,275, 17.6, 0.64; 4,4'-diaminodiphenylamine, 525, 44,850, 36.55, 1.36; 4,4'-diaminodiphenylmethane, 497, 34,500, 22.45, 0.834; benzidine, 527.5, 40,350, 39.0, 1.42; 3,3'-dichlorobenzidine, 520, 47,000, 33.4, 1.33; benzidine-3,3'-disulfonic acid, 515, 94,000, 25.0, 1.17; benzidine-2,2'-disulfonic acid, 502.5, 78,600, 10.54, 0.474; benzidine sulfone, 540, 32,930, 19.9, 0.787; benzidine sulfone-3,3'-disulfonic acid, 540, 55,850, 14.31, 0.712; diamino-2,2'-stilbenedisulfonic acid, 532, 82,800, 42.1, 2.015; p-phenylenediamine (monoazo deriv.), 510, 15,610, 25.25, 0.96; pphenylenediamine (bisazo deriv.), 515, 43,300, 32.2, 1.05; p,p'diaminodiphenylurea, 491, 40,200, 33.15, 1.302; 2,2'-dinitro-4,4'diaminodiphenyl-methane, 497, 27,350, 12.2, 0.508; .omicron.-aminophenol, 500, 25,600, 20.6, 0.785; 3,3'-diamino-4,4'-dihydroxydiphenyl-methane, 495, 42,600, 26.75, 1.035; 3-aminosalicylic acid, 505, 34,500, 13.5, 0.604; 3,3'-diamino-5,5'-methylenedisalicylic acid, 502, 64,300, 18.55, 0.84; The dyestuffs were prepd. by coupling the diazotized main component with 6-amino-1-naphthol-3sulfonic acid. Substantivities were assigned numerical values and were detd. spectrophotometrically, titration with Ti salts, and colorimetrically. The following general rules were proposed for a bisazo dyestuff to be substantive: (1) the mol. wt. must be fairly high, (2) at least 2 auxochromes must be linked by a long chain of conjugated double bonds (at least 8), (3) free rotation of aromatic nuclei must be possible (thus dyestuffs from benzidine-2,2'-disulfonic acid are acid dyestuffs which dye wool), (4) usually the dyestuff should not be a deriv. of a p,p'-diamine, (5) neg. substituents decrease the substantivity.

80-76-2, 4,6-Dibenzothiophenedisulfonic acid, 3,7-diamino-, IT 5,5-dioxide

(azo dyes from)

RN 80-76-2 CAPLUS

CN 4,6-Dibenzothiophenedisulfonic acid, 3,7-diamino-, 5,5-dioxide (8CI, 9CI) (CA INDEX NAME)

IT 858426-96-7, 4,6-Dibenzothiophenedisulfonic acid,
3-amino-7-(6-amino-1-hydroxy-3-sulfo-2-naphthylazo)-, 5,5-dioxide
(spectrum of)

RN 858426-96-7 CAPLUS

CN 4,6-Dibenzothiophenedisulfonic acid, 3-amino-7-(6-amino-1-hydroxy-3-sulfo-2-naphthylazo)-, 5,5-dioxide (5CI) (CA INDEX NAME)

L8 ANSWER 22 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1940:15386 CAPLUS Full-text

DOCUMENT NUMBER: 34:15386

ORIGINAL REFERENCE NO.: 34:2368g-i,2369a-i

TITLE: Dibenzofuran. XV. 1,4- and 1,4,6-Derivatives

AUTHOR(S): Gilman, Henry; Cheney, Lee C.

SOURCE: Journal of the American Chemical Society (1939), 61,

3149-56

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

Incidental to studies concerned with the bridging of the 1- and 9-positions in dibenzofuran types, series of 1,4- and 1,4,6-derivs. have been synthesized. In these compds. the 4- and 4,6-substituents are strong o,p-directors. 4-Hydroxy-6-methoxydibenzofuran (I) with HBr in AcOH gives 91.6% of 4,6dihydroxydibenzofuran (II), m. 200-2.degree.. II (0.115 mol), 0.345 mol of Me2SO4, 39 mL. of 60% KOH and Me2CO give a nearly quant. yield of 4,6dimethoxydibenzofuran (III), m. 128-9.degree.; picrate, deep yellow, m. 161-2.degree.. III and AcCl with AlCl3 in PhNO2 give 60% of the 1-Ac deriv. (IV), m. 178.5-9.5.degree... The oxime of IV, m. 203-4.degree., with PCl5 in C6H6 gives 76.4% of 1-acetamino-4,6- dimethoxydibenzofuran, m. 244-5.degree.; hydrolysis gives the HCl salt, m. 286-7.degree., of the 1-NH2 deriv. (V) of III, m. 162-2.5.degree.. I (4.28 g.) in 15 mL. 15% KOH and 385 mL. H2O, treated with the PhN2Cl from 0.02 mol PhNH2 at 3.degree., gives 56.6% of 1benzeneazo-4-hydroxy-6- methoxydibenzofuran, rust-colored, m. 175.degree.; Me2SO4 and KOH in Me2CO give 88% of the 4,6-di-MeO deriv., deep orange, m. 170.degree.; redn. gives 32.8% of V. 4-Methoxydibenzofuran (VI) and 10%

excess of (COCl)2 in PhNO2, treated with 10% excess AlCl3 at 0.degree. and allowed to stand at room temp. for 28 h., give a mixt. of 3 products; extn. of the amorphous product with 5% NaOH gives 4-methoxy-1-dibenzofurancarboxylic acid, m. 276-7.degree.; the alkali-insol. solid, extd. with AcOH, gives 18% of bis(4-methoxy-1-dibenzofuryl) ketone, m. 234.degree.; the AcOH-insol. portion is bi(4-methoxy-1-dibenzofuroyl), pale yellow, m. 329.degree., 34.6% yield. VI and ClCH2COCl with AlCl3 in PhNO2 give 53.2% of the 1-chloroacetyl deriv., m. 165-6.degree.; ClCOCO2Et gives 43% of the 1-ethoxalyl deriv., m. 113.degree.; hydrolysis with 15% NaOH gives 4-methoxy-1-dibenzofury1-.alpha.oxoacetic acid, pale yellow, m. 187.degree. (semicarbazone, m. 211.5-12.degree. (decompn.)). III (4.56 g.) and (COCl)2 with AlCl3 in PhNO2 give 0.37 g. 4,6-dimechoxy-1- dibenzofurancarboxylic acid, m. 297-8.degree.; 10.4% of bis(4,6-dimethoxy-1-benzofuryl) ketone, m. 254-5.degree., sol. in AcOH, and 60.7% of bi(4,6-dimethoxy-1-dibenzofuroyl), pale yellow, m. above 300.degree., insol. in AcOH. 3-Hydroxy-4-methoxydibenzofuran (VII) and HBr in AcOH give 88% of 3,4-dihydroxydibenzofuran (VIII), m. 164-4.5.degree.; di-Ac deriv., m. 104-5.degree.. VII and Me2SO4 with 10% NaOH give 81% of the 3,4-di-Me ether (IX), m. 60-1.degree.. IX and AcCl give 55.5% of the 1-Ac deriv. (X), m. 90.5-1.degree.. The oxime of X, m. 156-7.degree., is rearranged by PC15 in C6H6 to give 94% of 1-acetamino-3,4-dimethoxydibenzofuran, m. 156-7.degree.; hydrolysis gives 1-amino-3,4-dimethoxydibenzofuran, m. 162.5-3.degree., which also results in 10% yield on heating the 1-Br deriv. of IX with concd. NH4OH and CuBr for 14.5 h. at 220-30.degree.. III (22.8 g.) in 600 mL. AcOH and 100 mL. of M Br soln. in AcOH give 73% of the 1-Br deriv., m. 152.degree., and 12% of a product m. 144-7.degree.; III (3 g.) and 52.7 mL. of a 0.5 M Br-AcOH soln. give 74% of the 1,9-di-Br deriv. (XI), m. 167-8.degree.. II gives a nearly quant. yield of the 1,9-di-Br deriv., m. 239-40.degree. (decompn.); Me2SO4 gives XI. I gives 58.6% of the 1,3-di-Br deriv., m. 177-8.degree.; Me2SO4 gives the 1,3-di-Br deriv. of III, m. 173.5-4.degree.. IX forms 88.5% of the 1-Br deriv., m. 108.degree.; VII yields 54.6% of the 1-Br deriv., m. 161-2.degree., which was also prepd. from 1-bromo-3-amino-4methoxydibenzofuran through the diazo reaction in 21% yield. 4-Bromo-6methoxydibenzofuran (XII) and HI (d. 1.67) give 19% of the 6-HO analog, m. 138-9.degree.; FeCl3 gives a green color. XII, CuBr and NH4OH, heated in a steel bomb for 10 h. at 100.degree. and for 8 h. at 215.degree., give 51% of the HCl salt, m. 235-6.degree., of 4-amino-6-methoxydibenzofuran, m. 109.degree.; HBr in AcOH gives the 6-HO analog, m. 191.5-2.5.degree.. NaHSO3 and concd. NH4OH, heated at 185-95.degree. for 20 h., give 81% of 4,6diaminodibenzofuran, m. 152.degree.; HCl salt, m. 298.degree. (decompn.); picrate, red-brown, m. 213.degree. (decompn.); di-Ac deriv., m. 297-8.degree.. Di-Ac deriv. of II, m. 177.degree.. II and PhN2Cl give a dark brown compd., m. 228.degree. (decompn.), which is nearly pure 1,3,9-trisbenzeneazo deriv.; Me2SO4 gives 77% of 1,3,9-trisbenzeneazo-4,6-dimethoxydibenzofuran, redorange, m. 191-3.degree.. The 1-Ac deriv. of III, oxidized with I-KI in NaOHdioxane, gives 55.2% of 4,6-dimethoxy-1-dibenzofurancarboxylic acid (XIII), m. 297-8.degree.; this also resulted from carbonation of the Grignard reagent of the 1-Br deriv. of II; Me ester, m. 163.degree.. XIII gives an acid chloride, m. 147-50.degree.; CH2N2 gives 21.2% of diazomethyl 4,6-dimethoxy-1dibenzofuryl ketone, pale yellow, m. 151.degree. (decompn.); heating the ketone with concd. NH4OH and AgNO3 in dioxane gives 52% of the amide, m. 210-11.degree., of 4,6-dimethoxy-1- dibenzofurylacetic acid, m. 205.5-6.5.degree.. 3-Aminodibenzofuran, diazotized and reduced with SnCl2, gives 87.3% of the HCl salt, m. 242-3.degree., of 3-hydrazinodibenzofuran, pale yellow, m. 174-5.degree., which turns orange in the atm. 4-Aminodibenzofuran in abs. EtOH, reduced by Na in a N atm., gives 62% of 1,2,3,4-tetrahydro-6aminodibenzofuran, which is an oil at 0.degree.; HCl salt, pink, m. 228.degree. (decompn.); the diazo soln. with .beta.-C10H7OH gives a quant. yield of a brilliant carmine red dye, m. 199-201.degree.. 854394-18-6P, 4-Dibenzofuranol, 6-methoxy-1-phenylazo-

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1-Dibenzofuranamine, 4,6-dimethoxy-, -HCl 854395-26-9P,
1-Dibenzofuranamine, 4,6-dimethoxy- 854395-40-7P,
4,6-Dibenzofurandiol, 1,3,9-tris(phenylazo)- 854396-02-4P,
4-Dibenzofuranamine, 6-methoxy-, -HCl 854396-03-5P,
4-Dibenzofuranamine, 6-methoxy- 854396-65-9P, Dibenzofuran,
4,6-dimethoxy-1,3,9-tris(phenylazo)- 854396-67-1P, Dibenzofuran,
4,6-dimethoxy-1-phenylazo- 854397-18-5P, Dibenzofuran,
1-acetamido-4,6-dimethoxy-
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RL: PREP (Preparation)

(prepn. of)

RN 854394-18-6 CAPLUS

4-Dibenzofuranol, 6-methoxy-1-phenylazo- (4CI) (CA INDEX NAME) CN

RN 854394-30-2 CAPLUS CN 4-Dibenzofuranol, 6-amino- (4CI) (CA INDEX NAME)

854395-25-8 CAPLUS RN CN 1-Dibenzofuranamine, 4,6-dimethoxy-, -HCl (4CI) (CA INDEX NAME)

HC1

RN 854395-26-9 CAPLUS 1-Dibenzofuranamine, 4,6-dimethoxy- (4CI) (CA INDEX NAME) CN

RN 854395-40-7 CAPLUS

CN 4,6-Dibenzofurandiol, 1,3,9-tris(phenylazo)- (4CI) (CA INDEX NAME)

RN 854396-02-4 CAPLUS

CN 4-Dibenzofuranamine, 6-methoxy-, -HCl (4CI) (CA INDEX NAME)

HCl

RN 854396-03-5 CAPLUS

CN 4-Dibenzofuranamine, 6-methoxy- (4CI) (CA INDEX NAME)

RN 854396-65-9 CAPLUS

CN Dibenzofuran, 4,6-dimethoxy-1,3,9-tris(phenylazo)- (4CI) (CA INDEX NAME)

RN 854396-67-1 CAPLUS

CN Dibenzofuran, 4,6-dimethoxy-1-phenylazo- (4CI) (CA INDEX NAME)

RN 854397-18-5 CAPLUS

CN Dibenzofuran, 1-acetamido-4,6-dimethoxy- (4CI) (CA INDEX NAME)

L8 ANSWER 23 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

ORIGINAL REFERENCE NO.:

1939:29848 CAPLUS Full-text

DOCUMENT NUMBER:

33:29848 33:4238b-f

TITLE:

SOURCE:

Dibenzofuran. X. Aminohydroxy derivatives

AUTHOR (S):

Gilman, Henry; Jacoby, Arthur L.; Swislowsky, Jack Journal of the American Chemical Society (1939), 61,

954-6

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE:

Journal

LANGUAGE:

Unavailable

4-Acetaminodibenzofuran and HNO3 (d. 1.49) in Ac2O at -10.degree. give 35% of the Ac deriv. (I), pale yellow, m. 238.degree., of 3-nitro-4-aminodibenzofuran (II), deep yellow, m. 185-6.degree.. Catalytic redn. of I gives a quant. yield of 3-amino-4-acetaminodibenzofuran, silvery plates, m. 236-7.degree.; di-Ac deriv., m. 257.degree.. Redn. of II with Raney Ni and reaction with phenanthraquinone give dibenzo[a,c]benzofuro[2,3-h]phenazine, yellow, m. 277-8.degree.. 4-Hydroxydibenzofuran (III) and HNO3 in AcOH at -12.degree. give 25% of the 3-NO2 deriv. (IV), light yellow, m. 193.degree.; this also results from II through the diazo reaction. IV and CH2N2 give 65% of the 4-MeO deriv., yellow, m. 129.5.degree.. Nitration of III with concd. HNO3 in AcOH at 60.degree. gives 77% of the 3,8-di-NO2 deriv. (V), orange-red, m. 225.degree. (decompn.); this also results in a nearly quant. yield from IV.

V and CH2N2 give 83% of 3,8-dinitro-4-methoxydibenzofuran, orange, m.
177.degree.. The 2-isomer of III yields 80% of a yellow di-NO2 deriv., m.
240.degree. (decompn.), which is probably the 3,8-deriv. 4-Methoxydibenzofuran and HNO3 in Ac2O at -15.degree. to -20.degree. give 18% of the 1-NO2 deriv., m. 155.degree.; 1-NH2 deriv., pale lavender, m. 104.degree., which also results from 1-bromo-4-methoxydibenzofuran and concd. NH4OH with CuBr at 230-40.degree.. 4-Ethoxydibenzofuran gives 28% of the 1-NO2 deriv., yellow, m. 135-5.5.degree.; 1-NH2 deriv., m. 91.degree. (Ac deriv., m. 218.5.degree.).
3-Aminodibenzofuran, EtI, Na2CO3 and H2O, refluxed 48 h., give 70% of 3-diethylaminodibenzofuran, m. 68.degree.

IT 854394-20-0P, 4-Dibenzofuranol, 3,8-dinitro- 854396-46-6P, Dibenzofuran, 4-methoxy-3,8-dinitro-

RL: PREP (Preparation)

(prepn. of)

RN 854394-20-0 CAPLUS

CN 4-Dibenzofuranol, 3,8-dinitro- (4CI) (CA INDEX NAME)

RN 854396-46-6 CAPLUS
CN Dibenzofuran, 4-methoxy-3,8-dinitro- (4CI) (CA INDEX NAME)

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                 WPIDS/WPINDEX/WPIX manual codes updated
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NEWS 13 DEC 14
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                 functionality
NEWS 14
        DEC 18
                 CA/CAplus pre-1967 chemical substance index entries enhanced
                 with preparation role
NEWS 15
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                 CA/CAplus patent kind codes updated
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        DEC 18
                 MARPAT to CA/CAplus accession number crossover limit increased
                 to 50,000
                 MEDLINE updated in preparation for 2007 reload
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        DEC 27
                 CA/CAplus enhanced with more pre-1907 records
                 CHEMLIST enhanced with New Zealand Inventory of Chemicals
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NEWS EXPRESS
             MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
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0.21 0.21

FULL ESTIMATED COST

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STRUCTURE FILE UPDATES: 9 JAN 2007 HIGHEST RN 917076-17-6
DICTIONARY FILE UPDATES: 9 JAN 2007 HIGHEST RN 917076-17-6

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH June 30, 2006

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REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/ONLINE/UG/regprops.html

Uploading C:\Documents and Settings\ychu\Desktop\Case\10821642\10821642.str

chain nodes :

15 16 17 18 19 20 22 23 24

ring nodes :

1 2 3 4 5 6 7 8 9 10 11 12 13

chain bonds :

10-15 13-19 15-16 15-17 16-18 19-20 22-23 22-24

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-9 7-8 7-10 8-9 8-13 10-11 11-12 12-13

exact/norm bonds :

5-7 6-9 8-9 10-15 13-19 15-16 15-17 16-18 19-20 22-23 22-24

normalized bonds :

1-2 1-6 2-3 3-4 4-5 5-6 7-8 7-10 8-13 10-11 11-12 12-13

G1:0,S

G2:C,Q

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom

11:Atom 12:Atom 13:Atom 15:CLASS 16:CLASS 17:CLASS 18:Atom 19:Atom 20:CLASS

22:CLASS 23:CLASS 24:CLASS 26:Atom

L1 STRUCTURE UPLOADED

=> d

L1 HAS NO ANSWERS

L1 STR

Structure attributes must be viewed using STN Express query preparation.

Uploading C:\Documents and Settings\ychu\Desktop\Case\10821642\10821642a.str

chain nodes :

15 16 17 18 19 20 22

ring nodes :

1 2 3 4 5 6 7 8 9 10 11 12 13

chain bonds :

10-15 13-19 15-16 15-17 16-18 19-20

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-9 7-8 7-10 8-9 8-13 10-11 11-12 12-13

exact/norm bonds :

5-7 6-9 8-9 10-15 13-19 15-16 15-17 16-18 19-20

normalized bonds :

1-2 1-6 2-3 3-4 4-5 5-6 7-8 7-10 8-13 10-11 11-12 12-13

G1:0,S

## Match level :

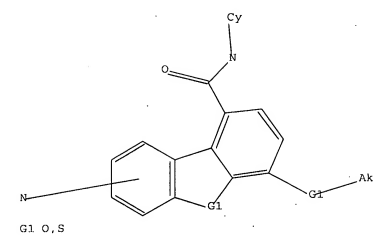
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 13:Atom 15:CLASS 16:CLASS 17:CLASS 18:Atom 19:Atom 20:CLASS 22:CLASS 23:Atom

L2 STRUCTURE UPLOADED

=> d

L2 HAS NO ANSWERS

L2 STR



Structure attributes must be viewed using STN Express query preparation.

=> s 12

SAMPLE SEARCH INITIATED 07:30:25 FILE 'REGISTRY'

SAMPLE SCREEN SEARCH COMPLETED - 131 TO ITERATE

100.0% PROCESSED 131 ITERATIONS 4 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*

BATCH \*\*COMPLETE\*\*

PROJECTED ITERATIONS: 1934 TO 3306

PROJECTED ANSWERS: 4 TO 200

L3 4 SEA SSS SAM L2

=> s 12 full

FULL SEARCH INITIATED 07:30:32 FILE 'REGISTRY'

FULL SCREEN SEARCH COMPLETED - 2760 TO ITERATE

100.0% PROCESSED 2760 ITERATIONS 69 ANSWERS

SEARCH TIME: 00.00.01

L4 69 SEA SSS FUL L2

=> file caplus

COST IN U.S. DOLLARS SINCE FILE TOTAL

ENTRY SESSION

FULL ESTIMATED COST 173.00 173.21

FILE 'CAPLUS' ENTERED AT 07:30:38 ON 11 JAN 2007
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FILE COVERS 1907 - 11 Jan 2007 VOL 146 ISS 3 FILE LAST UPDATED: 10 Jan 2007 (20070110/ED)

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## http://www.cas.org/infopolicy.html

=> s 14

L5 ·

=> d ibib abs hitstr tot

ANSWER 1 OF 4 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2006:361248 CAPLUS Full-text

DOCUMENT NUMBER:

144:412351

TITLE:

Process for the preparation of n-(3,5-dichloropyridin-

Not ODP.

Smethod for preparing

intermediate.

4-yl)-4-difluoromethoxy-8-methanesulfonamido-

dibenzo[b,d]furan-1-carboxamide

INVENTOR(S):

Gopalan, Balasubramanian; Gharat, Haxmikant Atmaram;

Chandrasekhar, Batchu; Karaunakaran, Usha; Pillai,

Bijukumar Gopinathan

PATENT ASSIGNEE(S):

SOURCE:

Glenmark Pharmaceuticals S.A., Switz.

PCT Int. Appl., 54 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PAT                                     | KIND DATE  |     |     |           | APPLICATION NO.             |     |      |            |                                   |       | DATE  |       |          |     |     |      |     |
|---|------------|-----|-----|-----------|-----------------------------|-----|------|------------|-----------------------------------|-------|-------|-------|----------|-----|-----|------|-----|
| 7                                       | 2006040652 |     |     |           | A2. 20060420<br>A3 20061026 |     |      |            | 1                                 | WO 2  | 005-  | IB30: | 20051012 |     |     |      |     |
|   |            | AE, | AG, | AL,       | AM,                         | AT, | AU,  | AZ,        | -                                 |       |       | •     |          |     |     |      |     |
|   |            |     |     | •         | •                           |     |      | DK,<br>IL, |                                   | •     |       |       |          |     |     |      |     |
|   |            |     |     |           |                             | •   |      | LV,        |                                   |       |       |       | 1        |     |     |      |     |
|   |            | •   | •   | -         | -                           |     | -    | PG,<br>TN, |                                   |       |       |       | 1        |     |     |      |     |
|   | RW.        |     | •   | ZM,<br>BG |                             | СХ  | CZ.  | DE,        | DK.                               | EE.   | ES.   | FT.   | FR       | GB. | GR. | ни.  | TE. |
|   |            | IS, | IT, | ·LT,      | LU,                         | LV, | MC,  | NL,        | PL,                               | PT,   | RO,   | SE,   | si¦,     | SK, | TR, | BF,  | ВJ, |
|   |            |     | •   |           |                             |     |      | GQ,<br>SD, |                                   |       |       |       | 1        |     |     |      |     |
| IIS                                     | 2006       | •   |     | MD,       |                             | -   |      | 0622       |                                   | IIS 2 | 005-1 | 2515  | .√<br>57 |     | 2   | 0051 | 013 |
| US 2006135779<br>PRIORITY APPLN. INFO.: |            |     |     |           | 77                          |     | 2000 | 0022       |                                   | US 2  | 004-  | 6184  | 74P      | 1   | P 2 | 0041 | 013 |
| . · ·                                   |            |     |     |           |                             |     |      |            | IN 2004-MU1099<br>US 2004-621981P |       |       |       | •        |     |     |      |     |

The present invention relates to a method of prepg. N-(3,5-dichloropyridin- 4-AB yl)-4-difluoromethoxy-8-methanesulfonamido-dibenzo[b,d]furan-1- carboxamide

and pharmaceutically acceptable salts thereof, such as its sodium salt, and novel intermediate compds. useful in the synthesis of the aforementioned compd. For example, reaction of 4-cyclopentyloxy-3- hydroxybenzaldehyde with 2-bromo-1-fluoro-4-nitrobenzene (70-77%), followed by cyclization, gave 4-cyclopentyloxy-8-nitro-1- formyldibenzofuran in 60-65% yield, which yielded the title compd. after 9 steps.

RN 685875-02-9 CAPLUS

CN

1-Dibenzofurancarboxamide, N-(3,5-dichloro-4-pyridinyl)-4-(difluoromethoxy)-8-nitro-(9CI). (CA INDEX NAME)

RN 685875-03-0 CAPLUS

CN 1-Dibenzofurancarboxamide, 8-amino-N-(3,5-dichloro-4-pyridinyl)-4-(difluoromethoxy)- (9CI) (CA INDEX NAME)

RN 778576-62-8 CAPLUS

CN

1-Dibenzofurancarboxamide, N-(3,5-dichloro-4-pyridinyl)-4-(difluoromethoxy)-8-[(methylsulfonyl)amino]- (9CI) (CA INDEX NAME)

IT 778576-63-9P

RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP (Preparation)

(prepn. of n-(3,5-dichloropyridin-4-yl)-4-difluoromethoxy-8-

methanesulfonamido-dibenzo[b,d]furan-1-carboxamide)

RN 778576-63-9 CAPLUS

CN 1-Dibenzofurancarboxamide, N-(3,5-dichloro-4-pyridinyl)-4(difluoromethoxy)-8-[(methylsulfonyl)amino]-, monosodium salt (9CI) (CF
INDEX NAME)

$$F_2CH = 0$$
 $O$ 
 $NH = S = Me$ 
 $C1$ 
 $NH$ 
 $C1$ 

Na

L5. ANSWER 2 OF 4 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2005:99226 CAPLUS Full-text

DOCUMENT NUMBER:

142:197859

TITLE:

Preparation of dibenzo[b,f]furan-1-carboxamides,

9H-carbazole-4-carboxamides, and dibenzo[b,d]thiophene-4-carboxamides as PDE4 inhibitors for the treatment of

Corrent app.

inflammatory and allergic disorders

INVENTOR(S):

Gopalan, Balasubramanian; Gharat, Laxmikant A.;

Lakdawala, Aftab D.; Karunakaran, Usha

PATENT ASSIGNEE(S):

Glenmark Pharmaceuticals, Inc. USA, USA

SOURCE:

U.S. Pat. Appl. Publ., 59 pp., Cont.-in-part of Appl.

No. PCT/IB04/000355.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 2
PATENT INFORMATION:

OTHER SOURCE(S):

GI

| PAT      | PATENT NO.    |                            |      |     |     | KIND DATE |      |      | APPLICATION NO.           |      |       |      |     |     | DATE       |     |     |    |  |
|----------|---------------|----------------------------|------|-----|-----|-----------|------|------|---------------------------|------|-------|------|-----|-----|------------|-----|-----|----|--|
|          |               |                            |      |     |     | -         |      |      |                           |      |       |      |     |     |            |     |     |    |  |
| US       | US 2005027129 |                            |      |     |     |           | 2005 |      |                           |      |       |      |     |     | 20040409   |     |     |    |  |
| WO       | WO 2004089940 |                            |      |     |     |           | 2004 | 1021 | WO 2004-18355             |      |       |      |     |     | 20040211   |     |     |    |  |
|          | W:            | AE,                        | AG,  | AL, | AM, | AT,       | AU,  | ΑZ,  | BA,                       | BB,  | BG,   | BR,  | BW, | BY, | BZ,        | CA, | CH, |    |  |
|          |               | CN,                        | CO,  | CR, | CU, | CZ,       | DE,  | DK,  | DM,                       | DZ,  | EC,   | EE,  | EG, | ES, | FI,        | GB, | GD, |    |  |
|          |               | GE,                        | GH,  | GM, | HR, | HU,       | ID,  | IL,  | IN,                       | IS,  | JP,   | KE,  | KG, | ΚP, | KR,        | KZ, | LC, |    |  |
|          |               | LK,                        | LR,  | LS, | LT, | LU,       | LV,  | MA,  | MD,                       | MG,  | MK,   | MN,  | MW, | MX, | MZ,        | NA, | NI, |    |  |
|          |               | NO,                        | NZ,  | OM, | PG, | PH,       | PL,  | PT,  | RO,                       | RU,  | SC,   | SD,  | SE, | SG, | SK,        | SL, | SY, |    |  |
|          |               | TJ,                        | TM,. | TN, | TR, | TT,       | TZ,  | UA,  | UG,                       | US,  | UZ,   | VC,  | VN, | YU, | ZA,        | ZM, | ZW  |    |  |
|          | RW:           | BW,                        | GH,  | GM, | KE, | LS,       | MW,  | MZ,  | SD,                       | SL,  | SZ,   | TZ,  | UG, | ZM, | ZW,        | AM, | ΑZ, |    |  |
|          |               | BY,                        | KG,  | KZ, | MD, | RU,       | TJ,  | TM,  | AT,                       | BE,  | BG,   | CH,  | CY, | CZ, | DE,        | DK, | EE, |    |  |
|          |               | ES,                        | FI,  | FR, | GB, | GR,       | HU,  | IE,  | IT,                       | LU,  | MC,   | NL,  | PT, | RO, | SE,        | SI, | SK, |    |  |
|          |               | TR,                        | BF,  | ВJ, | CF, | CG,       | CI,  | CM,  | GA,                       | GN,  | GQ,   | GW,  | ML, | MR, | ΝĒ,        | SN, | TD, | TG |  |
| PRIORITY | APP           | LN.                        | INFO | . : |     |           |      |      |                           | IN 2 | 003-1 | MU36 | 3   |     | A 20030411 |     |     |    |  |
|          |               | US 2003-519967P P 20031113 |      |     |     |           |      |      |                           |      |       |      |     | 113 |            |     |     |    |  |
|          |               |                            |      |     |     |           |      |      | WO 2004-IB355 A2 20040211 |      |       |      |     |     |            |     |     |    |  |

MARPAT 142:197859

ÒМе

Title heterocyclic tricycles I [wherein R1-R3, R5, R6, Ra = independently H, (un)substituted (cyclo)alkyl, (cyclo)alkenyl, alkynyl, (hetero)aryl, heterocyclyl(alkyl), etc.; R4 = NR5R6 (R5, R6 = H, alkyl, cycloalkyl, etc.), heterocyclyl; Ar = (un)substituted aryl(alkyl), heterocyclyl, heteroaryl; X = O, SOO-2, NRa; Y = CONR7, NR7SOO-2, SOO-2NR7, NR7CO; R7 = H, OH, ORa, (un)substituted alkyl, aryl, heterocyclyl; P = O, S; m = 0-3; n = 1-4; Ra = H, alkyl, cycloalkyl, etc.; and tautomers, regioisomers, stereoisomers, enantiomers, diastereomers, polymorphs, N-oxides, pharmaceutically acceptable salts, solvates, and compns. thereof] were prepd. as phosphodiesterase type 4 (PDE4) inhibitors. For example, N-(3,5-dichloropyrid-4-yl)-4-methoxy-8-aminodibenzo[b,f]furan-1- carboxamide (prepd. in six steps from isovanillin, 4-fluoronitrobenzene, and 4-amino-3,5-dichloropyridine) was coupled with methanesulfonyl chloride in THF and pyridine to give the sulfonamide II. The latter inhibited the PDE4-induced conversion of [3H] cAMP to the corresponding

TT

```
[3H] 5'-AMP with IC50 of 0.5058 nM. Thus, I and their pharmaceutical compns.
are useful for the treatment of immune disorders, inflammatory conditions,
allergic conditions, CNS diseases, and insulin resistant diabetes (no data).
778576-34-4P, N-(3,5-Dichloropyridin-4-yl)-4-methoxy-8-
[(methylsulfonyl)amino]dibenzo[b,d]furan-1-carboxamide
778576-37-7P, N-(3,5-Dichloropyridin-4-yl)-4-methoxy-8-
acetamidodibenzo [b,d] furan-1-carboxamide 778576-41-3P,
N-(3,5-Dichloropyridin-4-yl)-4-methoxy-8-[(hydroxycarbonylcarbonyl)amino]d
ibenzo[b,d]furan-1-carboxamide 778576-42-4P,
N-(3,5-Dichloropyridin-4-yl)-4-methoxy-8-[(ethoxycarbonylcarbonyl)amino]di
benzo[b,d]furan-1-carboxamide 778576-49-1P, N-(3,5-
Dichloropyridin-4-yl)-4-methoxy-8-[(phenoxycarbonyl)amino]dibenzo[b,d]fura
n-1-carboxamide 778576-54-8P, N-(3,5-Dichloropyridin-4-yl)-4-
methoxy-8-[[(N-methylpiperazin-4-yl)carbonyl]amino]dibenzo[b,d]furan-1-
carboxamide 778576-62-8P, N-(3,5-Dichloropyridin-4-yl)-4-
difluoromethoxy-8-[(methylsulfonyl)amino]dibenzo[b,d]furan-1-carboxamide
778576-66-2P, N-(3,5-Dichloropyridin-4-yl)-4-difluoromethoxy-8-
acetamidodibenzo [b, d] furan-1-carboxamide 778576-69-5P,
N-(3,5-Dichloropyridin-4-yl)-4-difluoromethoxy-8-
[(ethoxycarbonylcarbonyl)amino]dibenzo[b,d]furan-1-carboxamide
778576-70-8P, N-(3,5-Dichloropyridin-4-yl)-4-difluoromethoxy-8-
[(hydroxycarbonylcarbonyl)amino]dibenzo[b,d]furan-1-carboxamide
778576-72-0P, N-(3,5-Dichloropyridin-4-yl)-4-difluoromethoxy-8-
[[(fur-2-yl)carbonyl]amino]dibenzo[b,d]furan-1-carboxamide
778576-90-2P, N-(3,5-Dichloropyridin-4-yl)-4-methoxy-8-[[[(2-
ethoxy-2-oxoethyl)amino]carbonyl]amino]dibenzo[b,d]furan-1-carboxamide
778576-92-4P, N-(3,5-Dichloropyridin-4-yl)-4-methoxy-8-(2-ethoxy-2-
oxoethylamino)dibenzo[b,d]furan-1-carboxamide
RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic
preparation); THU (Therapeutic use); BIOL (Biological study); PREP
(Preparation); RACT (Reactant or reagent); USES (Uses)
   (PDE4 inhibitor; prepn. of tricyclic heterocycles as PDE4 inhibitors
   for treatment of immune and inflammatory disorders and insulin
   resistant diabetes)
778576-34-4 CAPLUS
1-Dibenzofurancarboxamide, N-(3,5-dichloro-4-pyridinyl)-4-methoxy-8-
```

[(methylsulfonyl)amino] - (9CI) (CA INDEX NAME)

TT

RN

CN

RN CN 778576-37-7 CAPLUS

1-Dibenzofurancarboxamide, 8-(acetylamino)-N-(3,5-dichloro-4-pyridinyl)-4-methoxy- (9CI) (CA INDEX NAME)

(prepn. of tricyclic heterocycles as PDE4 inhibitors for treatment of immune and inflammatory disorders and insulin resistant diabetes)

RN 836627-26-0 CAPLUS

> 1-Dibenzofurancarboxamide, N-(3,5-dichloro-1-oxido-4-pyridinyl)-4-(difluoromethoxy)-8-[(methylsulfonyl)amino]- (9CI) (CA INDEX NAME)

Current app.

L5 ANSWER 3 OF 4 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2004:878393 CAPLUS Full-text

DOCUMENT NUMBER:

141:366121

TITLE:

CN

Preparation of dibenzo[b,f]furan-1-carboxamides,

9H-carbazole-4-carboxamides, and dibenzo[b,d]thiophene-4-carboxamides as PDE4 inhibitors for the treatment of

inflammatory and allergic disorders

INVENTOR(S):

Gopalan, Balasubramanian; Gharat, Laxmikant Atmaram;

Lakdawala, Aftab Dawoodbhai; Karaunakaran, Usha

PATENT ASSIGNEE(S):

Glenmark Pharmaceuticals Ltd., India PCT Int. Appl., 121 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PAT           | CENT 1 | NO.  |     |     | KIND DATE   |     |      |      |     |      |      |      |     |        |     |       |     |    |
|---------------|--------|------|-----|-----|-------------|-----|------|------|-----|------|------|------|-----|--------|-----|-------|-----|----|
| WO 2004089940 |        |      |     |     | A1 20041021 |     |      |      |     | WO 2 |      |      |     | · //\. |     |       |     |    |
|               | W:     | ΑE,  | AG, | AL, | AM,         | AT, | AU,  | AZ,  | BA, | BB,  | BG,  | BR,  | BW, | BY,    | ΒZ, | CA,   | CH, |    |
|               |        | CN,  | CO, | ĊR, | CU,         | CZ, | DE,  | DK,  | DM, | DZ,  | EC,  | EE,  | EG, | ES,    | FI, | GB,   | GD, |    |
|               |        | GE,  | GH, | GM, | HR,         | HU, | ID,  | IL,  | IN, | IS,  | JP,  | KE,  | KG, | KP,    | KR, | ΚZ,   | LC, |    |
|               |        | LK,  | LR, | LS, | LT,         | LU, | LV,  | MA,  | MD, | MG,  | MK,  | MN,  | MW, | MX,    | MZ, | NA,   | NI, |    |
|               |        | NO,  | NZ, | OM, | PG,         | PH, | ΡL,  | PT,  | RO, | RU,  | SC,  | SD,  | SE, | SG,    | SK, | SL,   | SY, |    |
|               |        | ТJ,  | TM, | TN, | TR,         | TT, | TZ,  | UA,  | UG, | US,  | UZ,  | VC,  | VN, | YU,    | ZA, | ZM,   | zw  |    |
|               | RW:    | BW,  | GH, | GM, | KE,         | LS, | MW,  | MZ,  | SD, | SL,  | SZ,  | TZ,  | UG, | ZM,    | ZW, | AM,   | ΑZ, |    |
|               |        | BY,  | KG, | ΚZ, | MD,         | RU, | TJ,  | TM,  | AT, | BE,  | BG,  | CH,  | CY, | CZ,    | DE, | DK,   | EE, |    |
|               |        | ES,  | FI, | FR, | GB,         | GR, | HU,  | IE,  | IT, | LU,  | MC,  | NL,  | PT, | RO,    | SE, | SI,   | SK, |    |
|               |        | TR,  | BF, | ВJ, | CF,         | CG, | CI,  | CM,  | GA, | GN,  | GQ,  | GW,  | ML, | MR,    | NE, | SN,   | TD, | TG |
| AU            | 2004   | 2284 | 53  |     | A1          |     | 2004 | 1021 |     | AU 2 | 004- | 2284 | 53  |        | 2   | 0040  | 211 |    |
| CA            | 2522   | 023  |     |     | A1          |     | 2004 | 1021 | 1   | CA 2 | 004- | 2522 | 023 |        | 2   | 00402 | 211 |    |
| EP            | 1620   | 429  | •   |     | A1          |     | 2006 | 0201 |     | EP 2 | 004- | 7100 | 93  |        | 2   | 00402 | 211 |    |
|               | R:     | AT,  | BE, | CH, | DE,         | DK, | ES,  | FR,  | GB, | GR,  | IT,  | LI,  | LU, | NL,    | SE, | MC,   | PT, |    |
|               |        | IE,  | SI, | LT, | LV,         | FI, | RO,  | MK,  | CY, | AL,  | TR,  | BG;  | CZ, | EE,    | HU, | SK    |     |    |

| BR 2004009747          | Α  | 20060509 | BR | 2004-9747     |   | 20040211 |
|------------------------|----|----------|----|---------------|---|----------|
| CN 1829711             | A  | 20060906 | CN | 2004-80016048 |   | 20040211 |
| JP 2006522789          | T  | 20061005 | JP | 2006-506259   |   | 20040211 |
| US 2005027129          | A1 | 20050203 | US | 2004-821642   |   | 20040409 |
| NO 2005005316          | A  | 20060111 | NO | 2005-5316     |   | 20051110 |
| PRIORITY APPLN. INFO.: |    |          | IN | 2003-MU363    | Α | 20030411 |
|                        |    |          | US | 2003-519967P  | P | 20031113 |
|                        |    |          | WO | 2004-IB355    | W | 20040211 |

OTHER SOURCE(S):

CASREACT 141:366121; MARPAT 141:366121

GI

$$(R^3)$$
 m  $(R^4)$  n  $(R^4$ 

AB Title heterocyclic tricycles I [wherein R1-R3, R5, R6, Ra = independently H, (un) substituted (cyclo) alkyl, (cyclo) alkenyl, alkynyl, (hetero) aryl, heterocyclyl(alkyl), etc.; R4 = NR5R6, heterocyclyl; Ar = (un)substituted aryl(alkyl), heterocyclyl, heteroaryl; X = O, SOO-2, NRa; Y = CONR7, NR7SOO-2, SO0-2NR7, NR7CO; R7 = H, OH, ORa, (un) substituted alkyl, aryl, heterocyclyl; P = 0, S; m = 0-3; n = 1-4; and tautomers, regioisomers, stereoisomers, enantiomers, diastereomers, polymorphs, N-oxides, pharmaceutically acceptable salts, solvates, and compns. thereof] were prepd. as phosphodiesterase type 4 (PDE4) inhibitors. For example, N-(3,5-dichloropyrid-4-yl)-4-methoxy-8aminodibenzo[b,f]furan-1- carboxamide (prepd. in six steps from isovanillin, 4-fluoronitrobenzene, and 4-amino-3,5-dichloropyridine) was coupled with methanesulfonyl chloride in THF and pyridine to give the sulfonamide II. latter inhibited the PDE4-induced conversion of [3H] cAMP to the corresponding [3H] 5'-AMP with IC50 of 0.5058 nM. Thus, I and their pharmaceutical compns. are useful for the treatment of immune disorders, inflammatory conditions, allergic conditions, CNS diseases, and insulin resistant diabetes (no data). IT 778576-34-4P, N-(3,5-Dichloropyridin-4-yl)-4-methoxy-8-[(methylsulfonyl)amino]dibenzo[b,d]furan-1-carboxamide 778576-37-7P, N-(3,5-Dichloropyridin-4-yl)-4-methoxy-8acetamidodibenzo[b,d]furan-1-carboxamide 778576-41-3P, N-(3,5-Dichloropyridin-4-yl)-4-methoxy-8-[(hydroxycarbonylcarbonyl)amino]d ibenzo[b,d]furan-1-carboxamide 778576-42-4P, N-(3,5-Dichloropyridin-4-yl)-4-methoxy-8-[(ethoxycarbonylcarbonyl)amino]di benzo[b,d] furan-1-carboxamide 778576-49-1P, N-(3,5-Dichloropyridin-4-yl)-4-methoxy-8-[(phenoxycarbonyl)amino]dibenzo[b,d]fura n-1-carboxamide 778576-54-8P, N-(3,5-Dichloropyridin-4-yl)-4methoxy-8-[[(N-methylpiperazin-4-yl)carbonyl]amino]dibenzo[b,d]furan-1carboxamide 778576-62-8P, N-(3,5-Dichloropyridin-4-yl)-4-

778576-86-6 CAPLUS RN

1-Dibenzofurancarboxamide, 8-amino-4-methoxy-N-3-pyridinyl- (9CI) CN

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 6

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

CAPLUS COPYRIGHT 2007 ACS on STN ANSWER 4 OF 4

ACCESSION NUMBER:

2004:370918 CAPLUS Full-text

DOCUMENT NUMBER:

140:391192

TITLE:

Preparation of dibenzofuran/dibenzothiophene

derivatives useful for the treatment of inflammatory

and allergic disorders

INVENTOR(S):

Balasubramanian, Gopalan; Gharat, Laxmikant Atmaram;

Lakdawala, Aftab Dawoodbhai; Anupindi, Raghu Ram

PATENT ASSIGNEE(S):

SOURCE:

Glenmark Pharmaceuticals Ltd., India

PCT Int. Appl., 254 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO.    |    |     |     |     | KIN         | D   | DATE |     |      | APPLICATION NO. |          |     |     |     |      | DATE |      |  |  |
|---------------|----|-----|-----|-----|-------------|-----|------|-----|------|-----------------|----------|-----|-----|-----|------|------|------|--|--|
|               |    |     |     |     |             |     |      |     |      |                 |          |     |     |     |      |      |      |  |  |
| WO 2004037805 |    |     |     |     | A1 20040506 |     |      |     | WO 2 | 003-            | 20031008 |     |     |     |      |      |      |  |  |
|               | W: | ΑE, | AG, | AL, | AM,         | AT, | AU,  | AZ, | BA,  | BB,             | BG,      | BR, | BY, | BZ, | CA,  | CH,  | CN,  |  |  |
|               |    | CO, | CR, | CU, | CZ,         | DE, | DK,  | DM, | DZ,  | EC,             | EE,      | EG, | ES, | FI, | GB,  | GD,  | GE,  |  |  |
|               |    | CH  | CM  | HВ  | пH          | TD  | TI.  | TN. | TS   | σT.             | KE       | KG  | KР  | KB  | K 7. | LC.  | T.K. |  |  |

R4->NRSRA V present app.

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             OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM,
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PRIORITY APPLN. INFO.:
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                                              WO 2003-IB4442
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OTHER SOURCE(S):

MARPAT -140:391192

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Title compds. I [R1-3 = H, alk(en/yn)yl, cycloalkyl, etc.; P = O, S; n = 0-4; Ar = (un)substituted aryl, etc.; Y = carboxamido, aminosulfonyl, etc.] are prepd. For instance, 4-methoxydibenzofuran-1-carboxylic acid (prepn. given) is converted to the corresponding acid chloride (PhH, SOC12, reflux, 4 h) and treated with 4-amino-3,5-dichloropyridine (DMF/THF, NaH, -10.degree.) to give II. II has IC50 = 0.8 nM for PDE4. I are useful for the treatment of inflammatory conditions, diseases of the central nervous and insulin resistant diabetes.

IT 685874-79-7P, N-(3,5-Dichloropyridin-4-yl)-4-methoxy-8nitrodibenzofuran-1-carboxamide 685875-02-9P,
N-(3,5-Dichloropyridin-4-yl)-4-difluoromethoxy-8-nitrodibenzofuran-1carboxamide 685875-03-0P, N-(3,5-Dichloropyridin-4-yl)-4difluoromethoxy-8-aminodibenzofuran-1-carboxamide
RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic
preparation); THU (Therapeutic use); BIOL (Biological study); PREP
(Preparation); RACT (Reactant or reagent); USES (Uses)

(prepn. of dibenzofuran/dibenzothiophene derivs. useful for treatment of inflammatory and allergic disorders)

RN 685874-79-7 CAPLUS

CN 1-Dibenzofurancarboxamide, N-(3,5-dichloro-4-pyridinyl)-4-methoxy-8-nitro-(9CI) (CA INDEX NAME)